The Science of Chocolate
2nd Edition
Preface

In 1988 I wrote a paper for School Science Review, in which I described some of the science involved in chocolate making and followed this by two experiments that could be tried in the classroom. As a result of this I received letters from both pupils and teachers requesting more information or new experiments to try. Subsequently I was contacted by Chris Butlin, who was then developing a food option for the Salters' Physics Advanced Level course. This resulted in some of the science of chocolate being included in this option. The numerous talks given by my colleagues and myself to junior schools, societies and universities also convinced me that there was a genuine interest in this topic and that people were not just coming for the free samples.

When, therefore, the Royal Society of Chemistry asked me if I would write a full book on this topic, aimed at schools and universities, I agreed to do so, without realising the amount of work involved. It was very gratifying, however, to learn in 2006 that the sales were such that a second edition of the book was required. Several people had commented that they wished to know more about the health and nutrition aspects of chocolate and, as a lot of research has recently been carried out in the area, this seemed the ideal subject for a new chapter. In addition, in 2005 New Scientist published a book entitled Does Anything Eat Wasps? in which the question was asked about how are bubbles put into Aero®. Possible solutions are given here as part of a second new chapter.

Having graduated in physics, the book is naturally biased in this direction, although I have tried to include a substantial amount of chemistry and even some mathematics in the project work. Several
of the chemical terms used in industry are different from those taught in schools. I have attempted to use the current terminology and have included a glossary in the hope that it will be useful, if someone is not familiar with the term in the text. This glossary also explains some of the industry’s own technical names.

This book should be especially useful for someone studying food science at university or who is about to join the confectionery industry. Although a scientific background is required to understand the more difficult sections, such as fat chemistry or the Maillard reaction, most of the rest of the book should be readable by 16–18 year olds. Here I have attempted to show how concepts such as latent heat, relative humidity etc. play an important part in the making of something as apparently simple as chocolate. I hope that this in fact might prove to be a “painless” way of learning about them.

Several sections are relatively simple and can be adapted by teachers of GCSE science or even younger pupils. This is especially true of the projects described in Chapter 12. These are meant to be just basic ideas that can be adapted according to age. All use apparatus or ingredients that should be easy to make or obtain. The appropriate safety precautions must, of course, be taken especially for those involving glass, heat or chemicals.

Finally I would like to thank my wife Dorothy for her help with the book and our sons Christopher and Richard for their help with the diagrams, together with John Birkett, Patrick Couzens, Peter Geary, Duane Mellor and Lynda O’Neill for correcting the script, or testing the projects to ensure that they worked. I am also grateful to Awema, Blackwell Science, Loders Croklaan and Palsgaard Industri A/S for their permission to reproduce picture, diagrams and tables. In particular, Figures 1.2, 2.3, 3.5, 3.6, 3.10, 3.13, 3.14, 4.9, 4.11, 4.12, 5.2, 5.3, 5.8, 5.10, 5.13, 6.8, 7.1, 7.5 and 9.8 are all reproduced from *Industrial Chocolate Manufacture and Use* with the permission of Blackwell Science and Figures 1.3 and 1.4 with the permission of the Nestlé Archives, Vevey, Switzerland.

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Contents

Chapter 1  The History of Chocolate

<table>
<thead>
<tr>
<th>Section</th>
<th>Title</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.1</td>
<td>Chocolate as a Drink</td>
<td>1</td>
</tr>
<tr>
<td>1.2</td>
<td>Eating Chocolate</td>
<td>3</td>
</tr>
<tr>
<td>1.2.1</td>
<td>Chocolate Crumb</td>
<td>7</td>
</tr>
<tr>
<td>1.2.2</td>
<td>White Chocolate</td>
<td>8</td>
</tr>
<tr>
<td>1.3</td>
<td>Chocolate Marketing in the UK</td>
<td>9</td>
</tr>
<tr>
<td>1.4</td>
<td>Chocolate is Good for You</td>
<td>9</td>
</tr>
<tr>
<td></td>
<td>References</td>
<td>10</td>
</tr>
</tbody>
</table>

Chapter 2  Chocolate Ingredients

<table>
<thead>
<tr>
<th>Section</th>
<th>Title</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.1</td>
<td>Cocoa Beans</td>
<td>11</td>
</tr>
<tr>
<td>2.1.1</td>
<td>Cocoa Trees</td>
<td>11</td>
</tr>
<tr>
<td>2.1.2</td>
<td>Commercial Cocoa-Producing Countries</td>
<td>12</td>
</tr>
<tr>
<td>2.1.3</td>
<td>Cocoa Pods</td>
<td>14</td>
</tr>
<tr>
<td>2.1.4</td>
<td>Fermentation</td>
<td>15</td>
</tr>
<tr>
<td>2.1.4.1</td>
<td>Fermentation Procedure</td>
<td>15</td>
</tr>
<tr>
<td>2.1.4.2</td>
<td>Microbial and Chemical Changes</td>
<td>18</td>
</tr>
<tr>
<td>2.1.5</td>
<td>Drying</td>
<td>20</td>
</tr>
<tr>
<td>2.1.6</td>
<td>Storage and Transport</td>
<td>22</td>
</tr>
<tr>
<td>2.2</td>
<td>Sugar and Sugar Substitutes</td>
<td>23</td>
</tr>
<tr>
<td>2.2.1</td>
<td>Sugar and its Production</td>
<td>23</td>
</tr>
<tr>
<td>2.2.2</td>
<td>Crystalline and Amorphous Sugar</td>
<td>24</td>
</tr>
<tr>
<td>2.2.3</td>
<td>Lactose</td>
<td>26</td>
</tr>
<tr>
<td>2.2.4</td>
<td>Glucose and Fructose</td>
<td>28</td>
</tr>
<tr>
<td>2.2.5</td>
<td>Sugar Alcohols</td>
<td>28</td>
</tr>
</tbody>
</table>
Chapter 3 Cocoa Bean Processing

3.1 Bean Cleaning 39
3.2 Roasting and Winnowing 40
   3.2.1 The Problem of Bean Size Variation 40
   3.2.2 Winnowing 43
   3.2.3 Bean Roasting 44
   3.2.4 Nib and Liquor Roasting 44
   3.2.5 Roasters 45
   3.2.6 Chemical Changes during Roasting 46
   3.2.7 Maillard Reaction 47
3.3 Grinding Cocoa Nib 49
   3.3.1 Cocoa Mills 51
      3.3.1.1 Impact Mills 52
      3.3.1.2 Disc Mills 52
      3.3.1.3 Ball Mills 53
3.4 Cocoa Butter and Cocoa Powder Production 54
   3.4.1 Alkalising (Dutching) 54
   3.4.2 Cocoa Butter 55
   3.4.3 Cocoa Powder 57
References 57

Chapter 4 Liquid Chocolate Making

4.1 Chocolate Milling 61
   4.1.1 Separate Ingredient Grinding Mills 62
   4.1.2 Combined Milling 64
4.2 Chocolate Conching 68
   4.2.1 Chemical Changes 69
Contents

4.2.2 Physical Changes 70  
4.2.3 Viscosity Reduction 71  
4.2.4 Conching Machines 73  
   4.2.4.1 The Long Conche 73  
   4.2.4.2 Rotary Conches 74  
   4.2.4.3 Continuous Low Volume Machines 76  
4.2.5 The Three Stages of Conching 78  
References 79

Chapter 5 Controlling the Flow Properties of Liquid Chocolate

5.1 Viscosity 81  
5.2 Particle Size 84  
   5.2.1 Particle Size Distribution Data 84  
   5.2.2 Effect of Particle Size on Viscosity 86  
5.3 Effect of Fat Additions on Viscosity 90  
5.4 Moisture and Chocolate Flow 92  
5.5 Emulsifiers and Chocolate Viscosity 93  
   5.5.1 Lecithin 94  
   5.5.2 Polyglycerol Polyricinoleate 98  
   5.5.3 Other Emulsifiers 99  
5.6 Degree of Mixing 99  
References 101

Chapter 6 Crystallising the Fat in Chocolate

6.1 Structure of Cocoa Butter 104  
6.2 Different Crystalline Forms 107  
6.3 Pre-Crystallisation or Tempering 110  
6.4 Mixing Different Fats (Fat Eutectics) 112  
6.5 Chocolate Fat Bloom 116  
6.6 Some Types of Non-Cocoa Vegetable Fat 118  
   6.6.1 Cocoa Butter Equivalents 119  
   6.6.2 Enzyme Interesterification 120  
   6.6.3 Lauric Fat Cocoa Butter Replacers 121  
   6.6.4 Non-Lauric Fat Cocoa Butter Replacers 122  
   6.6.5 Low Calorie Fats 124  
References 124
Chapter 7  Manufacturing Chocolate Products

7.1  Tempering 125
    7.1.1  Liquid Chocolate Storage 125
    7.1.2  Tempering Machines 126
    7.1.3  Hand Tempering 128
    7.1.4  Temper Measurement 128

7.2  Moulding 132
    7.2.1  Solid Tablets 132
    7.2.2  Chocolate Shells 135

7.3  Enrobers 139
    7.3.1  Maintaining Tempered Chocolate 142

7.4  Solidifying the Chocolate 143
    7.4.1  Coolers 145

7.5  Panning 146
    7.5.1  Chocolate Coating 147
    7.5.2  Sugar Panning 150

References 152

Chapter 8  Analytical Techniques

8.1  Particle Size Measurement 153
8.2  Moisture Determination 156
8.3  Fat Content Measurement 158
8.4  Viscosity Determination 159
    8.4.1  Simple Factory Techniques 159
    8.4.2  The Standard Method 161

8.5  Flavour 163
8.6  Texture Monitoring 165
8.7  Crystallisation Amount and Type 167
    8.7.1  Nuclear Magnetic Resonance 167
    8.7.2  Differential Scanning Calorimetry 169

References 170

Chapter 9  Different Chocolate Products

9.1  Special Recipes 171
    9.1.1  Ice-cream coatings 172

9.2  Shape-Retaining Chocolate 173
Chapter 10 Legislation, Shelf Life and Packaging

10.1 Legislation 184
10.2 Shelf Life 186
10.3 Packaging 187
  10.3.1 Foil and Paper Wrap 189
  10.3.2 Flow Wrap 190
  10.3.3 Biopolymers 193
  10.3.4 Robotic Packing 194

References 195

Chapter 11 Nutrition and Health

11.1 Nutrition 196
  11.1.1 Fats 197
  11.1.2 Carbohydrates 198
  11.1.3 Proteins 199
11.2 Obesity 200
11.3 Tooth Decay 201
  11.3.1 Anti-Caries Factor in Cocoa 202
  11.3.2 Tooth-Friendly Milk Proteins 202
  11.3.3 Oxalic Acid 203
  11.3.4 Oral Clearance 203
11.4 Other Alleged Negative Reactions 203
  11.4.1 Migraine and Headaches 203
  11.4.2 Acne 203
  11.4.3 Allergies 204
Chapter 12 Experiments with Chocolate and Chocolate Products

Project 1: Amorphous and Crystalline Sugar 209
Project 2: Particle Separation 210
Project 3: Fat Migration 212
Project 4: Cocoa Butter Separation 213
Project 5: Chocolate Viscosity 214
Project 6: Particle Size of Chocolate 216
Project 7: Effect of Lecithin 217
Project 8: Changing the Continuous Phase 218
Project 9: Chocolate Temper 219
Project 10: Hardness Measurement 220
Project 11: Chocolate Composition and Product Weight Control 222
Project 12: Distributions and Probabilities 223
Project 13: Chromatography of Colours 223
Project 14: Effectiveness of Different Packaging Materials 225
Project 15: Viscosity and Flavour 227
Project 16: Heat-Resistance Testing 228
Project 17: Coefficient of Expansion 229
Project 18: The Maillard Reaction 231

Glossary 232

Subject Index 234
CHAPTER 1

The History of Chocolate

Chocolate is almost unique as a food in that it is solid at normal room temperatures yet melts easily within the mouth. This is because the main fat in it, which is called cocoa butter, is essentially solid at temperatures below 25 °C when it holds all the solid sugar and cocoa particles together. This fat is, however, almost entirely liquid at body temperature, enabling the particles to flow past one another, so the chocolate becomes a smooth liquid when it is heated in the mouth. Chocolate also has a sweet taste that is attractive to most people.

Strangely chocolate began as a rather astringent, fatty and unpleasant tasting drink and the fact that it was developed at all, is one of the mysteries of history.

1.1 CHOCOLATE AS A DRINK

The first known cocoa plantations were established by the Maya in the lowlands of south Yucatan about 600 AD. Cocoa trees were being grown by the Aztecs of Mexico and the Incas of Peru when the Europeans discovered central America. The beans were highly prized and used as money as well as to produce a drink known as chocolatl. The beans were roasted in earthenware pots and crushed between stones, sometimes using decorated heated tables and mill stones, similar to those illustrated in Figure 1.1. They could then be kneaded into cakes, which could be added to cold water to make a drink. Vanilla, spices or honey were often added and the drink whipped to make it frothy. The Aztec Emperor Montezeuma was said to have drunk 50 jars of this beverage per day.

Christopher Columbus bought back some cocoa beans to Europe as a curiosity, but it was only after the Spaniards conquered
Mexico that Don Cortez introduced the drink to Spain in the 1520s. Here sugar was added to overcome some of the bitter, astringent flavours, but the drink remained virtually unknown in the rest of Europe for almost a hundred years, coming to Italy in 1606 and France in 1657. It was very expensive and, being a drink for the aristocracy, its spread was often through connections between powerful families. For example, the Spanish princess Anna of Austria introduced it to her husband King Louis XIII of France and the French court in about 1615. Here Cardinal Richelieu enjoyed it both as a drink and to aid his digestion. Its flavour was not liked by everyone and one Pope in fact declared that it could be drunk during a fast, because its taste was so bad.

The first chocolate drinking was established in London in 1657 and it was mentioned in Pepys’ Diary of 1664 where he wrote that “jocolatte” was “very good”. In 1727 milk was being added to the drink. This invention is generally attributed to Nicholas Sanders. During the eighteenth century, White’s Chocolate House became the fashionable place for young Londoners, while politicians of the day went to the Cocoa Tree Chocolate House. These were much less rowdy than the taverns of the period. It remained however, very much a drink for the wealthy.

One problem with the chocolate drink was that it was very fatty. Over half of the cocoa bean is made up of cocoa butter. This will melt in hot water making the cocoa particles hard to disperse as well as looking unpleasant, because of fat coming to the surface. The Dutch, however, found a way of improving the drink by
removing part of this fat. In 1828 Van Houten developed the cocoa press. This was quite remarkable, as his entire factory was manually operated at the time. The cocoa bean cotyledons (known as cocoa nibs) were pressed to produce a hard “cake” with about half the fat removed. This was milled into a powder, which could be used to produce a much less fatty drink. In order to make this powder disperse better in the hot water or milk, the Dutch treated the cocoa beans during the roasting process with an alkali liquid. This has subsequently become known as the Dutching process. By changing the type of alkalising agent, it also became possible to adjust the colour of the cocoa powder.

1.2 EATING CHOCOLATE

Having used the presses to remove some of the cocoa butter, the cocoa powder producers were left trying to find a market for this fat. This was solved by confectioners finding that “eating” chocolate could be produced by adding it to a milled mixture of sugar and cocoa nibs. (The ingredients used to make dark chocolate are shown in Figure 1.2.) If only the sugar and cocoa nibs were milled and mixed together they would produce a hard crumbly material. Adding the extra fat enabled all the solid particles to be coated with

![Figure 1.2](image-url)  
**Figure 1.2** Unmilled ingredients used to make dark chocolate: A, sugar; B, cocoa butter; C, cocoa nibs. 


fat and thus form the hard uniform bar that we know today, which will melt smoothly in the mouth.

Almost twenty years after the invention of the press in 1847, the first British factory to produce a plain eating chocolate was established in Bristol in the UK by Joseph Fry.

Unlike Van Houten, Fry used the recently developed steam engines to power his factory. It is interesting to note that many of the early chocolate companies, including Cadbury, Rowntree and Hershey (in the USA) were founded by Quakers or people of similar religious beliefs. This may have been because their pacifist and teetotal beliefs prevented them from working in many industries. The chocolate industry was, however, regarded as being beneficial to people. Both Cadbury and Rowntree moved to the outside of their cities at the end of the 1990s, where they built “garden” villages for some of their workers. Fry remained mainly in the middle of Bristol and did not expand as quickly as the other two companies. It eventually became part of Cadbury.

With the development of eating chocolate the demand for cocoa greatly increased. Initially much of the cocoa came from the Americas, with the first cocoa plantation in Bahia in Brazil being established in 1746. Even earlier, however, the Spaniards took cocoa trees to Fernando Po (Biyogo), off the coast of Africa, and this soon became an important growing area. In 1879 a West African blacksmith took some plants home to the Gold Coast (now Ghana). The British governor realised its potential and encouraged the planting of trees, with the result that Ghana has become a major source of quality cocoa. Other European powers also encouraged the growing of cocoa in their tropical colonies, e.g. France in the Ivory Coast (Côte d’Ivoire), which is now the world’s largest producer of cocoa.

The chocolate made by Fry was initially a plain block and it was only in 1875 that the first milk chocolate was made by Daniel Peter in Switzerland. Chocolate cannot contain much moisture, because water reacts with the sugar and turns melted chocolate into a paste rather than a smoothly flowing liquid (see Project 5 in Chapter 12). As little as 2% moisture can give a product a poor shelf life as well as an inferior texture. This meant that Daniel Peter had to find some way of drying the plentiful supply of liquid milk that he found in his own country. He was helped in this by the recent development of a condensed milk formula by Henri Nestlé. This
meant that he had much less water to evaporate, and he was able to remove the remaining amount using relatively cheap water-powered machines. In most countries milk chocolate products are now much more popular than plain chocolate ones. In the early 1900s Daniel Peter was challenged to prove that he did in fact invent milk chocolate, so he took his original notebook to the lawyer to get it stamped. The original page together with the lawyer’s mark is reproduced in Figure 1.3.
At this time most of the milk chocolate-like bars were still used to make drinks. Figure 1.4 shows an early 1900s’ advertisement for Peter’s company. At the foot of it is a triangular bar which was known as Peter’s Delta chocolate. It was made so that it could easily be broken into smaller triangular pieces, each of which would dissolve in a cup full of hot water.

In order for the chocolate to feel smooth on the tongue when it melts in the mouth, the solid non-fat particles must be smaller than 30 microns (1000 microns = 1 mm). The chocolates made by Fry and Peter were ground using granite rollers, but still had a gritty texture. This was because of the presence of some large particles and some groups of particles joined together to form agglomerates, also because the fat was not coating the particles very well. In addition, the chocolate tended to taste bitter because of the presence of some acidic chemicals (see Chapter 4).

In 1880 Rodolphe Lindt, in his factory in Berne in Switzerland, invented a machine which produced a smoother, better tasting
chocolate. This machine was known as a conche, because its shape was similar to the shell with that name (Figure 1.5). It consisted of a granite trough, with a roller, normally constructed of the same material, which pushed the warm liquid chocolate backwards and forwards for several days. This broke up the agglomerates and some of the larger particles and coated them all with fat. At the same time moisture and some acidic chemicals were evaporated into the air, producing a smoother, less astringent tasting chocolate. A schematic diagram of the chocolate making processes is shown in Figure 1.6.

1.2.1 Chocolate Crumb

In the early part of the twentieth century the milk used to make chocolate had poor keeping qualities. This caused problems for the chocolate industry, whose major sales were at Christmas, a time of the year when there was a very limited supply of fresh milk. In the
UK and some other countries this lead to the development of an intermediate ingredient called “chocolate crumb”.

The cocoa nibs contain substances known as antioxidants (see Chapter 11). These restrict the breaking up of the fats, which would normally make the milk fat turn sour. In addition, sugar was known to extend the shelf life of foods (it is used in jams etc.). The chocolate manufacturers therefore added sugar and cocoa to the milk and dried them together. This produced chocolate crumb, which had a shelf life of at least a year. Milk produced during the spring peak could then be used to make chocolate the following Christmas. The drying process, however, introduced some cooked flavours into the chocolate and it is for this reason that many UK milk chocolates taste different from some continental European ones, which are made from milk powder.

### 1.2.2 White Chocolate

The first white chocolate was made in 1930. It was made from sugar, milk powder and cocoa butter. The preserving qualities of
the cocoa antioxidants are mainly in the dark cocoa material. This means that white chocolate does not keep as well as milk chocolate, and also that it should be kept in a non-transparent wrapper, as light will speed up the decomposition of the milk fat.

1.3 CHOCOLATE MARKETING IN THE UK

As the technology improved, chocolate was used to coat other ingredients, or to be part of a product rather than just a bar. In the 1930s many of these were developed and have remained popular to this day. Good examples of this are KitKat®, Mars Bar® and Smarties®. At this time products also became known under brand name rather than that of the manufacturer. Some companies, like Cadbury tended to give both equal prominence.

During the war few cocoa beans were shipped from the plantations and strict rationing was introduced. Many leading brands were not produced at all. Rationing in the UK ended in April 1949, but the rush to buy was so great that, by June, 60% of confectionery shops had nothing left to sell. Rationing was reimposed until February 1953.

Consumption rose very quickly, but over the last 10 years has been more constant with an average of about 9 kg/person per annum of chocolate confectionery being eaten in many West European countries (this does not include chocolate biscuits). Germany is the highest at about 11 kg/person per annum. This makes the confectionery industry a very important one. The combined sales of sugar and chocolate confectionery in the UK are more than tea, newspapers and bread put together.4

1.4 CHOCOLATE IS GOOD FOR YOU

Antioxidants in food are known to protect the body against chemicals called free radicals that damage cells. Cocoa is a known source of antioxidants and in 1999 doctors from the National Institute of Public Health and the Environment in Bilthoven in The Netherlands examined chocolate for its catechins content. These are from the family of flavonoids, which are among the most powerful antioxidants. They found that dark chocolate contained 53.5 mg/100 g, which is four times that in tea. Drinking a cup of tea a day is said to reduce the chance of a heart attack.5 Since then a lot
of work has been carried out, which has shown the beneficial effects of cocoa with respect to heart disease and possibly some cancers (see Chapter 11).

In spite of its perceived negative image within the general public with respect to obesity, tooth decay, acne and migraine, the scientific evidence is that chocolate does not play a significant role in any of these, provided it is eaten in normal amounts as part of a balanced diet (Chapter 11).

REFERENCES

CHAPTER 2

Chocolate Ingredients

2.1 COCOA BEANS

2.1.1 Cocoa Trees

To be called “chocolate”, a product must contain cocoa. The cocoa or cacao tree (Theobroma cacao L.) originated in South and Central America, but is now grown commercially in suitable environments between 20° north and 20° south. These areas have a high average temperature ($\geq 27^\circ$C) throughout the year and a constant high humidity, arising from a plentiful rainfall (1500–2500 mm). The soil should be deep, rich and well drained, and normally be less than 700 m above sea level, as strong winds will damage the crop.

The trees are relatively small, 12–15 m in height, and grow naturally in the lower level of the evergreen rainforest. In commercial plantations they are often sheltered by intercropping trees such as coconut and banana. Its leaves are evergreen and are up to about 300 mm in length. Trees start bearing pods after 2–3 years, but it is 6 or 7 years before they give a full yield.

There are four types of cocoa. Criollo has beans with white cotyledons and a mild flavour. The trees are, however, relatively low yielding. Most cocoa is Forastero, which is more vigorous and often grown on smallholdings (a family’s cultivated land that is smaller than a farm) in West Africa. The third form, Trinitario, is usually thought to be a hybrid of the other two types. The fourth type is Nacional, which is grown only in Ecuador and probably originates from the Amazonian area of Ecuador. Nacional cocoa
produces beans with a full cocoa flavour with additional floral, spicy flavours.

The trees are attacked by many pests and diseases. Some of the most serious are:

- capsids (insects that feed on sap, causing damage to plant tissue)
- black pod disease (fungi which attack mainly the pods, making them rot)
- witches’ broom disease (fungal attack causing growths or brooms to develop from leaf buds, but also affects flowers and pods)
- cocoa pod borer moth (larva bores into the pods and affects the development of the beans).

### 2.1.2 Commercial Cocoa-Producing Countries

There are three major cocoa growing regions: West Africa, South-East Asia and South America. These are shown in Figure 2.1. The cocoa supplies from individual countries has changed dramatically over the last twenty years due to economic changes as well as pests and diseases.

![Figure 2.1](image)

**Figure 2.1** Map showing the major cocoa growing countries of the world.
Bahia in Brazil was a major cocoa growing area producing over 400 000 tonnes in the mid-1980s. It now produces less than half of this, largely due to destruction by witches’ broom disease. Instead of being a major exporter of beans, almost all are now used for internal consumption.

About 40% of the world’s crop (1.4 million tonnes) is now grown in the Ivory Coast (Côte d’Ivoire). Production has increased dramatically over the last 30 years and a large proportion of European chocolate it made from this source of beans. Most is grown on smallholdings. Recent political instabilities, however, may make future supplies less certain.

Ghana is the second highest producer with around 20% of the world’s crop – 700 000 tonnes. It also has a good reputation for producing quality beans and is the standard against which other bulk beans are measured. Increasing amounts of cocoa are being processed locally.

Nigeria produces some cocoa, but the establishment of the oil and other industries resulted in alternative employment and a reduction in cocoa production. This has recently stabilised at about 200 000 tonnes, but many of the trees are old.

Indonesia has expanded its cocoa growing and is now producing almost as much as Ghana. The flavour of the cocoa depends not only on the cocoa type, e.g. whether it is Criollo or Forastero, but also on the climate and soil conditions etc. For some specialist chocolates, normally dark ones, beans are obtained from specific areas. These fine or flavour cocoas, often Criollo, are produced in many smaller growing areas such as Ecuador, the Caribbean Islands and Papua New Guinea.

Malaysia built up a fairly big cocoa production in the 1980s, but this has declined rapidly, partly due to pod borer infestation and also due to the greater profitability of other crops such as oil palm.

In addition to the flavour of the beans, the fat contained within it also changes according to the area of production. In general, the nearer the equator that the tree is grown the harder is the fat, i.e. melts at a higher temperature. This means that Malaysian cocoa butter is relatively hard, whereas most Brazilian cocoa butter is much softer. The first is better for chocolates that will be sold in summer, whereas the soft is preferable for frozen products, such as choc-ices, where the fat is hardened by the cold conditions. See Chapter 6 and Project 10 in Chapter 12.
2.1.3 Cocoa Pods

Tiny flowers (Figure 2.2), up to 100 000 in number per season, grow on the branches and trunk of the tree throughout the year. These grow into small, green pods called cherelles (Figure 2.3), but take 5–6 months to develop into mature pods (Figure 2.4) between 100 mm and 350 mm long. They weigh from 200 g to more than 1 kg and exist in a wide variety of shapes and colours depending on the variety. Each pod contains some 30–45 beans.

The pods are carefully cut off the tree with a machete (cutlass), where they are within reach. For the higher branches it is necessary to use a special knife attached to a long pole, as shown in Figure 2.5. Pods are normally harvested every 2–4 weeks over a period of several months, as they do not all ripen at the same time.

The pods are opened with a machete or cracked with a wooden club. The beans are oval in shape and covered in a white pulp (mucilage). The beans are separated from the majority of this pulp by hand.

The beans consist of an outer shell or testa surrounding two cotyledons (called nibs) and a small germ (the embryo plant). The cotyledons store the food for the developing seedling and also its first two leaves. Much of the food is in the form of a fat (cocoa

![Figure 2.2 Flowers on cocoa trees.](image)
butter) which accounts for over half the dry weight of the bean. The moisture content of the bean at this stage is about 65%.

2.1.4 Fermentation

2.1.4.1 Fermentation Procedure. Correct fermentation is essential to produce a good flavour in the final chocolate. It is a process in which the bean is killed, so that it cannot be spoiled by germination. In addition, certain chemicals are formed which on heating give the taste of cocoa, whereas these chemicals themselves taste completely different or may not even taste of anything at all. These are known as flavour precursors as they lead to the flavour, but aren’t it themselves. Unfermented beans may be pressed to produce cocoa butter, but the remaining solid cocoa material is not normally used to make chocolate.

A lot of cocoa trees are grown by smallholders and the method of fermentation is traditional, although in some countries there have been attempts to modernise it. There are two main types of method: heap fermentation and box fermentation.

In West Africa, heap fermentation is widely used. Between 25 and 2500 kg of fresh beans, together with a small amount of the
white pulp, are placed in a heap and then covered with banana leaves (Figure 2.6). The process normally lasts from 5–6 days, with the actual length being determined by experience. Some farmers turn the beans after 2 or 3 days. The smaller heaps often produce the better flavours.

The larger plantations, particularly in Asia use the box fermentation technique. The wooden boxes may hold between 1 tonne and 2 tonnes of beans, which are designed with outlet holes or slits, usually in the base (Figure 2.7). These provide ventilation and let
the water that comes out of the beans and pulp run away. These may be up to a metre deep, but shallower ones (250–500 mm) often give a better flavour due to the improved ventilation. The beans are tipped from one box to another each day to increase aeration and give a more uniform treatment. Usually the fermentation period is similar to that for the heap procedure, although some plantations may take a day or two longer.
2.1.4.2 Microbial and Chemical Changes. What actually happens during fermentation has been the subject of much research and is still not entirely clear. As the bean shell remains intact, it is not possible for micro-organisms to react directly with the cotyledons inside, which are the part that is used to make chocolate. So, in a way, this is not a true fermentation process at all.

During the fermentation, however, the temperature rises dramatically during the early stages and three days of heat are thought to be sufficient to kill the bean. Following its death, enzymes (a type of catalyst capable of greatly increasing the rate of breakdown of substances like fat into simpler components, but which comes out of the reaction unchanged itself) are released. These cause the rapid decomposition of the beans’ food reserves and form sugars and acids, which are the precursors of the chocolate flavour that was described earlier.
The process is, however, much more complicated than this, as a more usual fermentation process takes place outside the bean. Here there is some of the white pulp, which is very sugar-rich and able to react with the yeasts that are also present to form acids and ethanol, much in the same way as occurs during brewing. This ethanol activates other bacteria, e.g. ethanoic (acetic) acid and lactic acid bacteria, which then convert it into their respective acids. The ethanol and acids are able to pass through the shell into the bean. This change in acidity (pH) hastens the death of the bean.

As was noted earlier, there are different ways of fermenting and these too will give rise to different flavours. For example, in box fermentation the beans are moved every day. This aerates the beans....
and stimulates those bacteria that require oxygen (e.g. *Acetobacter*) and encourages the production of ethanoic acid. Other reactions, for instance those involving yeasts, are retarded by the presence of oxygen, so less ethanol is formed. This means that cocoas that are box fermented are more likely to taste acidic than the same type of cocoa that has been heap fermented. In order to overcome some of this acidity, some box fermentation processes shorten the fermentation time and reduce the number of turnings.

Many other important reactions also occur. The proteins and peptides react with polyphenols to give the brown colour associated with cocoa, whilst other flavour precursors are formed by reactions between sucrose and proteins. Of particular importance is the formation of amino acids. Proteins are composed of a series of amino acids joined alternatively by the acid and amine (NH$_2$) components. Amino acids can be represented as:

\[
\begin{array}{c}
  \text{H}_2\text{N} \quad \text{C} \quad \text{COOH} \\
  \text{Z} \\
  \text{H}
\end{array}
\]

2.1

where Z = H, CH$_3$, C$_2$H$_5$, CH$_2$OH etc.

(In practice the amino acid is ionic and either the NH$_2$ is protonated or the COOH is dissociated depending on the pH of the surroundings.)

Many of the proteins break up during the fermentation into these acids. Twenty forms exist, including valine and glycine, which are very important in chocolate flavour formation.

Further details of chemical changes during fermentation are given by Fowler$^1$ and Dimick and Hoskin.$^2$

2.1.5 Drying

Following fermentation, the beans must be dried before they can be transported to the chocolate making factories. Failure to do this will result in moulds growing on the beans. These give the chocolate a strong, nasty flavour and so cannot be used. Beans must also not be over-dried. Those with a moisture content of less than 6% become very brittle, which makes subsequent handling and processing much more difficult.
Where the weather permits, the beans are usually sun dried. They are spread out during the day in layers about 100 mm thick on mat, trays or terraces. They are raked at intervals, and heaped up and protected at night or when it rains. In Central and South America a roof on wheels is used to cover the beans, which are laid on the floor. In Ghana split bamboo mats are placed on low, wooden tables. The mats can be rolled up when it rains. Here it normally takes about a week for the beans to dry to the required 7–8% moisture level, which is too low for moulds to grow. In other areas beans are dried on moveable tables that can be put under cover when necessary (Figure 2.8). A major problem of sun drying is the risk of contamination from the surroundings and from farm and wild animals wandering amongst the beans. This means that precautions must be taken in handling them when they reach the chocolate making factory (see Chapter 3).

In other countries, particularly in Asia, the weather may be too wet and artificial drying is required. Sometimes wooden fires are lit in a chamber below the drying area, and the hot gas is led through a flue beneath the drying platform and then out through a vertical chimney. A major problem here is that of smoke leaking from the flue. This, like mould, gives the beans an unpleasant, harsh flavour.
and prevents them from being used for chocolate making. Forced-air dryers are better, as are efficient heat exchangers that stop smoky contaminants reaching the beans. If the drying is too quick, the beans will taste very acidic and it is better to dry them at lower temperatures or intermittently over a longer period.

2.1.6 Storage and Transport

The beans must be stored so that they do not pick up water as they will become mouldy once their moisture level rises above 8%. Traditionally they are stored in 60–65 kg jute sacks (Figure 2.9). These are strong, stackable and allow the moisture to pass through. They are also biodegradable. As chocolate is a very delicate flavour, the beans must also be stored well away from other goods such as spices, which might result in off-flavours in the chocolate.

The beans are often transported in the holds of ships. At the point of loading the temperature will be about 30 °C, but very soon the temperature in the North Atlantic will be nearer freezing point. If the beans have a moisture content of 8%, their equilibrium relative humidity is about 75%. In other words, if the relative humidity is below 75%, the beans will dry out and they will pick up moisture at

Figure 2.9  Cocoa in jute sacks ready for loading onto a ship.
higher humidities. The moisture level must not exceed 8%, so beans must never be stored at higher humidities. In the ships, however, this is difficult. The humidity is already high when the sacks are loaded and the drop in temperature causes a rapid rise in the relative humidity to 100% (the dew point). Moisture will condense on the ship’s structure and sometimes get into the sacks, making the beans mouldy. The sacks should therefore never be in contact with the cold surfaces and the absorbent mats placed on top of them. The ship’s hold should also be ventilated to remove the moist air.

2.2 SUGAR AND SUGAR SUBSTITUTES

Traditionally chocolate has been made containing about 50% sugar, mostly in the form of sucrose, but with some lactose from the milk components in milk chocolate. Diabetics are unable to eat much sugar, so other recipes were initially developed to incorporate fructose (a different form of sugar, also found in honey) or non-sugar bulk sweeteners such as sorbitol. More recently there has been a requirement for lower calorie or “tooth friendly” chocolates, so other sugar substitutes have been developed.

2.2.1 Sugar and its Production

Sucrose (also known as saccharose) is produced from both sugar beet and sugar cane. Both give the same natural crystalline disaccharide material. It is called a disaccharide because it is composed of two single sugars (monosaccharides) chemically linked together. These sugars, called glucose and fructose, are present in equal proportion and can be separated by acidic treatment or by using an enzyme called invertase. The resulting mixture of the two sugars is called invert sugar.

\[
\text{sugar} \quad \text{C}_{12}\text{H}_{22}\text{O}_{11} + \text{H}_2\text{O} \rightarrow \text{invert sugar} \quad 2\text{C}_6\text{H}_{12}\text{O}_6 \quad \text{fructose + glucose (isomers)}
\]  

Lactose is also a disaccharide and is made up of a combination of glucose and galactose. Many of the sugar substitutes, like sorbitol, are sugar alcohols.

Sugar beet contains about 14–17% sucrose. The beet is cleaned and sliced, and the sugar, together with some mineral and organic
impurities, are washed out with warm water. Slaked lime is added to precipitate out these impurities and carbon dioxide is then bubbled through the solution. This precipitates out the excess slaked lime as calcium carbonate. The solids are removed by filtration to give a 15% sugar solution, which is the evaporated to 65–70%. Vacuum evaporation and centrifugation are then used to purify and crystallise the sucrose. It is not possible to recover all the sugar in a single processing stage and white sugar requires three or four different steps.

Sugar cane has a sucrose content of 11–17%. The raw juice is squeezed from the crushed stalks, often using roller mills. The remaining material can be used to make paper, cardboard or hardboard etc. This juice contains more invert sugar than was the case with beet sugar. This makes it more difficult to crystallise and a gentler treatment is required to get rid of the impurities, otherwise an undesirable brown colour would form. Alternative treatments involve lower temperatures or the use of sulphur dioxide. Hydrocyclones and bow-shaped sieves are used to clarify the liquid, and crystallisation procedures are similar to those used for beet sugar.

2.2.2 Crystalline and Amorphous Sugar

Crystal sugar is extremely pure, normally being more than 99.9% and rarely less than 99.7%. It can be purchased with different crystal sizes, which are approximately as follows.

- Coarse sugar 1.0–2.5 mm grain size
- Medium–fine sugar 0.6–1.0 mm grain size
- Fine sugar 0.1–0.6 mm grain size
- Icing sugar 0.005–0.1 mm grain size

Most chocolate manufacturers use medium–fine sugar, although some ask for a defined particle size spectra, as this may aid the flow of the final chocolate (see Chapter 5).

All these sugars are in a crystalline form. Sucrose crystals can in fact be grown to be several centimetres long and can take many crystalline forms. All are however birefringent. That means that if the crystals are placed in a polarising microscope, with the polaroid filters set so that no light is transmitted, they bend the light and are seen as bright images (Figure 2.10).
Sugar can also exist as a glass, *i.e.* a non-crystalline, though solid, structure. A good example is a clear, boiled-sugar sweet, which is often mint flavoured. This happens when sucrose solutions are dried too quickly and the individual molecules do not have time to form the crystalline structure when the water is removed. One way to manufacture amorphous sugar is to freeze-dry a sucrose solution. Amorphous sugar is not birefringent as it does not possess a structure such that it can bend the light in a polarising microscope. There are other ways of determining amorphous sugar in sucrose systems (see Project 1 in Chapter 12).

Amorphous sugar is important in chocolate making as it can effect both the flavour and the flow properties of liquid chocolate. Its surface is very reactive and can easily absorb any flavours that are nearby. It is also formed from crystalline sucrose at high temperatures. These may occur when sugar is milled. If there is no other material around, the sugar may take up a metallic note. (This can be demonstrated by finely grinding sugar in a food mixer with a metal blade or bowl and then dissolving the sugar in water; it will taste metallic compared with a solution made from the original material.) On the other hand, if it is milled together with cocoa,
some of the volatile cocoa flavours are absorbed by the amorphous sugar rather than escape into the atmosphere as they would otherwise do. This will then produce a more intense flavour chocolate. Care must be taken when milling sugar, especially by itself, because of the high risk of an explosion.

The amorphous state is an unstable one, and in the presence of water it will turn into crystalline material. Once the change has taken place the moisture is expelled, as crystalline sucrose is essentially anhydrous. About half the mass of chocolate is sucrose, so the particles within it are very close together. The moisture on the surface makes them stick together. This eventually builds up a skeleton, which holds the sugar together even if the fat melts and runs out of it. This is the basis of a method used to create a chocolate suitable for sale in hot climates. If the chocolate has not yet been solidified, the stickiness on the surface of some of the sugar greatly increases the viscosity of the liquid chocolate.

Crystalline sugar can also absorb moisture, depending on its surrounding conditions. The storage conditions that should be used can be determined by means of sorption isotherms. Figure 2.11 illustrates the curve for sugar at 20 °C. As was noted earlier, the equilibrium relative humidity is the relative humidity at which water is neither taken in nor given out. This means that between 20% and 60% humidity the sugar will maintain a moisture of between 0.01% and 0.02%. At higher humidities the moisture content increases dramatically. Damp sugar may be microbiologically contaminated. In addition it will stick together and form lumps, even if the humidity is reduced again.

In the chocolate industry, sugar is stored in large silos containing many tonnes. Great care must be taken with the storage conditions as otherwise the silo will block up and fail to empty. Very often the air inside them is dehumidified.

2.2.3 Lactose

Like sucrose, lactose is a disaccharide, this time consisting of monosaccharides called glucose and galactose. It is part of cows’ milk (see next section) and is therefore found in all milk chocolates. It is sometimes used as crystalline lactose to replace part of the sucrose. As it is much less sweet than normal sugar, it reduces the overall sweetness of the chocolate. The crystalline form is a
monohydrate, i.e. it contains one molecule of water, which it does not loose even if the temperature is raised to 100 °C.

There are in fact two crystalline forms α- and β-lactose. The α form is produced by most conventional processes and is slightly less sweet and less soluble than the other form. Both are non-hygroscopic, i.e. they do not readily take up water. The amorphous form is however very hygroscopic.

When lactose is added in the form of spray dried milk powder it is normally in an amorphous form. This can be demonstrated by Project 1 in Chapter 12. This glassy state is able to hold in some of the milk fat and so make it unavailable to help the chocolate flow (Chapter 5).

At elevated temperatures lactose is able to take part in what is called the Maillard or browning reaction. This is the type of
reaction that occurs to produce cooked flavours when toasting food. It will be described in more detail in Chapter 3.

Some people are lactose intolerant, i.e. their body reacts against this type of sugar and they must limit the amount of milk products they eat.

### 2.2.4 Glucose and Fructose

Although these monosaccharides (single sugars) combine together to make the disaccharide sucrose, they are not normally used to make chocolate. Glucose, also called dextrose, crystallises as a monohydrate and is very difficult to dry completely. It normally contains some water and also rapidly absorbs it from the surrounding air (i.e. it is very hygroscopic). This moisture makes the liquid, melted chocolate very thick, because it tends to stick the sugar particles together.

Fructose is also very hygroscopic. It is found naturally in fruits and honey. It is sometimes found in special chocolates for diabetics, as, unlike sucrose, it does not raise the blood sugars when eaten. It does however need special processing conditions, especially with regard to temperature and humidity.

### 2.2.5 Sugar Alcohols

Sugar alcohols (polyols) are used to replace sucrose in chocolate, when it is required to make a lower calorie or a sugar-free product. Sucrose is normally regarded as containing 4.0 kcal g\(^{-1}\) (17 kJ g\(^{-1}\)). Although the different sugar alcohols probably have different caloric values, in Europe, for legislative purposes, they are regarded as having 2.4 kcal g\(^{-1}\) (10 kJ g\(^{-1}\)). Like fructose, they are suitable for diabetics, but, unlike it, they are suitable to make non-cariogenic chocolate – in other words, chocolates that are not damaging to the teeth. One of them, xylitol, which is found naturally in many mushrooms and fruits, cannot be fermented by most of the bacteria in the mouth and in some countries is regarded as beneficial to the teeth.

Other common sugar alcohols include sorbitol, mannoitol, isomalt and lactitol. Some require to be processed into chocolate at relatively low temperatures, to prevent them forming gritty lumps. All tend to have a laxative effect. The EU Scientific Committee on Foods recognised this to be important, but stated that a consumption of 20 g per day was unlikely to have a harmful effect.
There is also a big difference in sweetness between the different sugar alcohols. The relative values with respect to sucrose are given in Table 2.1. Intense sweeteners, such as aspartame, may be required with some polyols, such as sorbitol, to offset their lower sweetness.

When some substances dissolve in water, energy is required to enable this to take place. This occurs for sucrose (see Project 1 in Chapter 12) and results in a cooling of the water to enable the molecules to separate and dissolve. This takes place to a much greater degree with the sugar alcohols. Table 2.2 shows the relative degrees of cooling and is particularly noticeable for xylitol. This property is often regarded as being undesirable, as chocolate is not expected to have a cooling effect.

### 2.2.6 Polydextrose

Polydextrose is a polysaccharide (i.e. made up of several sugars), which is found in many low calorie chocolate bars. It has a legislative

<table>
<thead>
<tr>
<th>Table 2.1 Relative degree of sweetness of different sugars and sugar alcohols (Krüger).</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sugar</strong></td>
</tr>
<tr>
<td>-----------</td>
</tr>
<tr>
<td>Sucrose</td>
</tr>
<tr>
<td>Fructose</td>
</tr>
<tr>
<td>Glucose</td>
</tr>
<tr>
<td>Xylitol</td>
</tr>
<tr>
<td>Maltitol</td>
</tr>
<tr>
<td>Sorbitol</td>
</tr>
<tr>
<td>Mannitol</td>
</tr>
<tr>
<td>Isomalt</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 2.2 Relative cooling effect of bulk sweeteners (Krüger).</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sugar</strong></td>
</tr>
<tr>
<td>-----------</td>
</tr>
<tr>
<td>Sucrose</td>
</tr>
<tr>
<td>Polydextrose</td>
</tr>
<tr>
<td>Lactitol (anhydrous)</td>
</tr>
<tr>
<td>Isomalt</td>
</tr>
<tr>
<td>Maltitol</td>
</tr>
<tr>
<td>Fructose</td>
</tr>
<tr>
<td>Lactitol (monohydrate)</td>
</tr>
<tr>
<td>Sorbitol</td>
</tr>
<tr>
<td>Xylitol</td>
</tr>
</tbody>
</table>
calorific value in Europe of 1 kcal g⁻¹, *i.e.* less than one half of the value for the sugar alcohols, and it has much less of a laxative effect. It is made up from the monosaccharide sugar glucose with small amounts of the sugar alcohol sorbitol. Because it is amorphous, it reacts with water to give out heat (just like amorphous sucrose, see Project 1 in Chapter 12). This means that it gives a warming sensation when dissolved in the saliva in the mouth. It also dries out the mouth, sometimes making the chocolate harder to swallow.

### 2.3 MILK AND OTHER DAIRY COMPONENTS

In most countries of the world much more milk chocolate is bought and eaten than both dark and white put together. It tends to be softer than dark chocolate and has a creamier taste and texture.

The majority of cows’ milk is water, but, as was mentioned earlier, moisture destroys the flow properties of liquid chocolate so only the anhydrous components can be used. Typically these form about 13.5% of the liquid milk and their composition is illustrated in Figure 2.12. The largest component, at just under 5%, is lactose, the disaccharide sugar described in the previous section. There is almost the same amount of milk fat and about 3.5% of the protein.

![Milk Composition](image)

**Figure 2.12** Relative composition of dried milk.
Minerals account for about 0.7%. Calcium in particular is regarded as being very beneficial to health.

2.3.1 Milk Fat

This is the second largest component in dehydrated milk and is vital in giving milk chocolate its distinctive texture and flavour release. It also changes its snap and can inhibit the formation of the white powdery surface on chocolate, which is composed of large fat crystals and is known as fat “bloom” (see Chapter 6).

The more fat that is present in a liquid chocolate the easier it will flow, both when making sweets and in the mouth. It is also relatively expensive, so the manufacture needs to make the best use of the fat present.

Butter fat is almost entirely liquid at room temperature, so there is a limit to the amount that can be added to chocolate for it still to remain hard. To make matters worse there is a phenomena, called fat eutectics, which means that when two fats are added together the resultant mixture is often softer than would be expected. This is described in more detail in Chapter 6. This softness does however reduce the waxiness in the mouth if harder fats than cocoa butter are present. Fractionate milk fats are also available. These are made by separating out the higher or the lower melting point fractions. Certain ones are said to produce a harder chocolate than the normal milk fat, whereas others are thought to give better bloom prevention.

The fat is 98% triacylglycerols (triglycerides), i.e. three acids combined with a glycerol molecule. The remaining significant component is phosphoglyceride (phospholipid) (mainly lecithin) as well as diacylglycerols (diglycerides) (two acids combined with glycerol) and sterols. A typical fatty acid composition in milk is given in Table 2.3.

Milk fat, however, has a limited shelf life as it can be oxidised or attacked by enzymes (lipolysis). The enzymes accelerate the break up of the acids into shorter chain free acids, which have a rancid off-flavour and make the chocolate unacceptable. When this type of reaction occurs with cocoa butter, however, the acids formed are largely tasteless, so the chocolate remains acceptable.

The initial result of oxidation is the formation of peroxides (–O–O–). These have no taste themselves, but decompose to produce unpleasant off-flavours. A measurement of the amount
of peroxide present is used to detect the early stages of deterioration. In order to keep the milk fat for longer periods, contact with oxygen must be minimised. Sometimes the air in the packaging is replaced by nitrogen and an oxygen barrier packaging is used. Chilled storage is preferred and the presence of copper and iron must be avoided as these act as catalysts for the oxidation process.

### Table 2.3 Fatty acid composition of milk fat (Haylock and Dodds⁴).

<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>Weight (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C4:0* Butyric</td>
<td>4.1</td>
</tr>
<tr>
<td>C6:0 Caproic</td>
<td>2.4</td>
</tr>
<tr>
<td>C8:0 Caprylic</td>
<td>1.4</td>
</tr>
<tr>
<td>C10:0 Capric</td>
<td>2.9</td>
</tr>
<tr>
<td>C10:1 Decenoic</td>
<td>0.3</td>
</tr>
<tr>
<td>C12:0 Lauric</td>
<td>3.5</td>
</tr>
<tr>
<td>C14:0 Myristic</td>
<td>11.4</td>
</tr>
<tr>
<td>C16:0 Palmitic</td>
<td>23.2</td>
</tr>
<tr>
<td>C18:0 Stearic</td>
<td>12.4</td>
</tr>
<tr>
<td>C18:1 Oleic</td>
<td>25.2</td>
</tr>
<tr>
<td>C18:2 Linoleic</td>
<td>2.6</td>
</tr>
<tr>
<td>C18:3 Linolenic</td>
<td>0.9</td>
</tr>
<tr>
<td>Others</td>
<td>10.0</td>
</tr>
</tbody>
</table>

* The first number is the number of carbon atoms in the molecule and the second is the number of double bonds.

2.3.2 Milk Proteins

Not only do these add to the nutritional content of the chocolate, they are also important in determining its flavour, texture and liquid flow properties. A milk chocolate has a creaminess which depends to a large extent on the balance between these proteins and the more acidic, cookie flavour from the beans. If the protein proportion is reduced, the product becomes much less creamy. Also, like lactose, if the proteins are subjected to water and heat, they can take part in the Maillard (browning) reaction, which introduces cooked flavours into the chocolate.

There are two very different types of protein, namely caseins and whey proteins. There are four to five times as much caseins as there are whey proteins. Some of their properties are summarised in Table 2.4.

Caseins act as emulsifiers, *i.e.* as interfaces between two different media. In chocolate this is likely to be between the solid and fat
components (cf. lecithin in Chapter 5). Its actual role is not understood, but calcium caseinate has been shown to produce thinner chocolate, whereas whey proteins make the same chocolate much thicker. The water binding properties of the caseins are also beneficial to the chocolate flow.

Although helping the flavour of some chocolates, the casein flavour itself is not particularly pleasant and may be undesirable in other confectionery products.

### 2.3.3 Milk Powders

Milk can be dried to produce a wide range of different powders. Figure 2.13 shows a flow chart of the dairy processes used to make powders for chocolate making.

The most common powders used for chocolate making are skim milk and full cream milk powder. With the former, milk fat is added at the chocolate making stage so that both powders can be used to make chocolates with the same overall milk components content. They will, however, have different flavours, textures and liquid flow properties. This is in part due to different heat treatments during the drying, but also due to the different state of the fat. With skim milk and milk fat, all the fat is free to react with the particles and the cocoa butter, whereas many full cream milk powders have fat tightly bound within the individual particles. This means that there is less fat to help the flow or to soften the cocoa butter when full cream powders are used.

For many years the milk powders were dried on hot rollers. These machines were expensive and hard to maintain in a hygienic condition, so currently most milk powders are produced by spray drying. This involves converting the milk, which has been partly pre-dried to about 50% moisture, into a mist (“atomised”) so that it has a very large surface area. The droplets are exposed to a flow of hot air in the drying chamber. The air provides heat to evaporate
Figure 2.13  Flow chart of the dairy processes used to make powders for chocolate manufacture.
the water and also acts as a carrier for the powder, which is then collected using cyclone separators (where centrifugal forces remove particles from air rotating at high speeds c.f. bagless vacuum cleaners) or filter systems. This powder often consists of fine, hollow, spherical particles similar to the ones illustrated in Figure 2.14. This is very different from the more plate-like structure of the roller dried milk powder shown in Figure 2.15.

When the fat in a chocolate is liquid, these particles must pass by one another when the chocolate flows, e.g. when it moves around the mouth. The different shaped particles will flow past each other differently, which may result in a change of viscosity. The spherical particles also contain fat inside them, whereas the rollers press most of it onto the surface. This means that roller dried full cream milk powder chocolates are softer and flow more easily than spray dried ones. Modifications can be made to the spray drying procedure to make the particles more crystalline so as to enable them to release the fat more easily.

The milk may also be heat treated before spraying, to introduce a more cooked flavour. Sometimes extra fat is added to give a high-fat powder, so that the chocolate manufacturer has no need to make subsequent additions during the production stage.

![Figure 2.14 Electron microscope picture of spray dried skim milk powder.](image-url)
Other milk powders contain sugar and are a form of chocolate crumb.

2.3.4 Whey and Lactose Powders

Both these powders are used in the making of certain types of chocolate and flavoured coating. They are by-products of cheese and casein manufacture and have the advantage that they are less sweet than sugar and so can replace some of it when a reduction in sweetness is required.

The acidity (pH) of the cheese making greatly affects the mineral content of the whey powder. Normally the more acidic the whey, the higher will be the mineral content. These may give rise to off-flavours in the chocolate and demineralised whey powders may be preferable.

If the protein is removed from the whey, lactose is left. This is then concentrated and crystallised.

2.4 CHOCOLATE CRUMB

As was noted in Chapter 1, chocolate crumb exploits the antioxidant properties in cocoa to produce a milk-containing chocolate.
ingredient with a long shelf life. It also gives a slightly cooked flavour to the chocolate.

There are many ways of producing crumb, but all require the final product to have a moisture content of between 0.8% and 1.5%. At this level the water activity (equilibrium relative humidity divided by 100) is so low that micro-organisms cannot grow. Care must be taken as to where it is stored, because, as with sugar, it will pick up water under moist surroundings.

Some manufacturers dissolve sugar in fresh or concentrated milk, whereas others add water to a mixture of sugar and milk powder. Multiple effect plate dryers may be used to dry this to 80–90% solids. Milled cocoa beans (cocoa liquor) are mixed with this material, which is then dried under vacuum in either a batch or a continuous process. At this stage there is heat, moisture, lactose and proteins. These are ideal conditions for the Maillard reaction to occur. This gives the crumb a brown colour and a caramel type flavour. This flavour is quite different from the one which would be obtained if sugar were heated alone. Also subsequent processes have almost no moisture present, so it is not possible to get this type of flavour during further manufacturing.

Most milk chocolates that are available on the market throughout the world are made from very similar ingredients, yet have very different flavours. Some manufacturers have specific “house flavours”, Cadbury and Hershey perhaps being the most well known. Very often these are due to specific changes during the crumb making process, where changes in holding times, acidity and temperature can give rise to very different flavours in the final chocolate. For this reason, chocolate crumb making conditions are often kept a closely guarded secret.

REFERENCES


**ADDITIONAL READING**

Traditionally cocoa beans were transported to the country where the chocolate was going to be manufactured, which was normally situated in a temperate climate. Increasingly, however, the cocoa growing countries are processing their own beans to produce cocoa liquor. This has the advantage that the liquor is much easier to transport – the problem of moisture affecting the beans on board ships was noted earlier. In addition, the shell, which is essentially a waste product, is not transported thousands of miles just to be thrown away. The disadvantage is that the manufacturer has less control over the times and temperature of the bean processing, which have an effect on the final chocolate flavour. One solution is to partially process the beans, with the final heat treatment being carried out at the chocolate factory. Wherever the process is carried out, it involves cleaning the beans, removing the shell and some form of roasting.

3.1 BEAN CLEANING

As many of the beans are dried on the ground, they often contain sand, stones, iron, plant material etc. These must be removed for two reasons. Firstly, many of these impurities are very hard and will damage the machinery which is used to grind the beans. Secondly, the organic contaminants will burn during the roasting process and give off gases likely to spoil the cocoa flavour. The cleaning is therefore carried out at the beginning of the chocolate making process.

Normally there are several different procedures, which are combined to remove the various types of waste material. Iron is removed by magnets, whereas dust can be drawn off by suction. Stones may be a similar size to the beans, but they are a different
density. They can be separated by vibrating them together on a grid, which is set at an angle to the horizontal. Air passes through the grid and blows the beans higher than the stones. As they are nearer the vibrating grid, the stones are moved towards the top, where they fall off into a collecting bag. The air transports the beans towards the lower part of the grid, from where they go on to the next stage of processing.

3.2 ROASTING AND WINNOWING

The cotyledons (nibs) must be roasted before they can be made into chocolate. This is to change the flavour precursors into the chemicals which actually taste of chocolate. In addition, the high temperature, coupled with the remaining moisture in the beans, will kill any microbiological contaminants, such as salmonellae, which may be present on the beans from when they were dried on the open ground.

Many manufacturers use a concept known as hazard analysis of critical control points (HACCP). This means that the whole chocolate making procedure is evaluated to determine any possible source of harm to the consumer. As the beans are purchased from bulk suppliers and may be contaminated with harmful bacteria etc., it is necessary to treat all the beans as if they were a potential hazard until the risk has been removed. The roasting procedure will do this, so once this has been carried out the cocoa is absolutely safe. Analysis of the beans at this stage, known as a critical control point, will confirm this.

The main risk is then that the contamination from the untreated beans will be transferred to the rest of the factory. For this reason, the pre-roasting procedures, such as cleaning, are normally carried out in a separate building. The operators are also obliged to change their clothing before entering the rest of the factory.

3.2.1 The Problem of Bean Size Variation

Cocoa beans come in a variety of sizes, depending on the country of origin, the climatic conditions, the season of the year the pod was picked, and numerous other factors. Traditionally the beans were roasted in small batches of perhaps a few hundred kilograms, often in ball-shaped roasters (see Figure 3.1). The operator could remove a few beans from the roaster or the attached cooling tray,
and from the smell of the beans was able to adjust the temperature and times to ensure that the correct flavour had been developed. Beans could also be segregated so that they were of a similar size within each batch. With modern factories, which need to process several tonnes of beans per hour, this is no longer possible.

The problem caused by having a variety of bean sizes is illustrated in Figure 3.2. Here it can be seen that when the roasting conditions are set for the average size of bean, the smaller ones are over-roasted, whereas as the centres of the larger beans are not roasted enough. This means that in the latter case not all the flavour precursors will have been converted, so the chocolate flavour will be low. In the case of the small bean, further compounds may have been produced, which are not necessarily of the required flavour.

The difficulty with the large beans is illustrated in Figure 3.3. Here the temperature is measured in the roaster itself and at several locations within a bean. Even after a relatively long time (15 minutes) the temperature of the centre of the bean fails to reach that of the outside. The difference in formation of the different flavour compounds can also be shown by carrying out analysis by systems such as high pressure liquid chromatography (HPLC, see Chapter 8) on different sizes of bean.
In order to overcome this, two alternative methods have been developed. In the first only the centre of the bean is roasted, so the pieces are much smaller, and the heat can more easily get to the middle. This process is known as nib roasting. In the second, the nib is finely ground it is turned into cocoa liquor (also known as mass).

Figure 3.2  Effect of bean size on degree of roasting.

Figure 3.3  Change of temperature in the roaster and within a bean during the early part of roasting.
Here the cocoa butter has been released from the cells within the bean, so that when it becomes warm it turns into a liquid. This liquid can then be heated in a process called liquor roasting. A schematic flow diagram of the different roasting methods is given in Figure 3.4.

Both these methods require the shell to be removed before the roasting process. This must be done carefully, because the shell often has silica particles attached, often originating from when the beans were dried on the ground. These, together with the shell, which is also very hard, will damage any grinding equipment during the chocolate making. In addition there are sometimes legal requirements, which state that only a small amount of shell can be present in anything that is sold as chocolate. The shell is also said to give an inferior flavour to the chocolate. The small amount of shell fat present is different from cocoa butter and, like milk fat (Chapters 2 and 6), has a softening effect on the chocolate.

### 3.2.2 Winnowing

Winnowing is the process of separating the shell and some of the germ from the rest of the bean. As its name suggests, it relies on the
same principles as are used to separate the corn from the chaff during grain harvesting.

It is desirable to keep the central cotyledons (nibs) in as large pieces as possible so that they can separate more easily from the shell. Any small pieces that stay with the shell will be thrown away with it, so economically it is very important to carry out winnowing correctly.

Any broken beans are initially separated, to prevent them being broken further, and go straight to separation process. The remaining beans are broken, often by sending them individually at high speed against impact plates. These, too, then go to the vibrating sieves.

The shell is largely fibrous material and is normally in the form of a flat platelet. The nib, on the other hand, is normally much more spherical and, being over half fat, is much denser. When the two are vibrated together the lighter shell will come to the top (cf. the separation of beans and stones). If air is drawn upwards through this mixture, the lighter shell with its larger surface area will rise, whereas the heavier nib falls for further processing. This principle is illustrated in Project 2 in Chapter 12.

3.2.3 Bean Roasting

This process is still used by many chocolate manufacturers. It has the major advantage that the roasting tends to help separate the shell from the nib. This makes breaking and winnowing relatively easy.

There are, however, two other disadvantages in addition to the problem of the different sized beans. When the heat is applied, the cocoa butter melts. Some of it is free enough to migrate into the shell, where it remains and is thrown away following winnowing. It is estimated that up to 0.5% of cocoa butter is lost in this way. In addition, extra energy is required to heat the nib through the shell. All the energy used in heating the shell is also totally wasted. It is estimate that up to 44% extra energy is needed compared with the other forms of roasting.

3.2.4 Nib and Liquor Roasting

Because the shell is relatively firmly attached to the cotyledons until the bean has been heated, some form of pre-treatment is normally
required before winnowing. This normally consists of exposing the beans very rapidly to a source of intense heat provided by saturated steam or infrared radiation. This heats the surface, but the centre remains much cooler and no flavour changing reactions occur. Water inside the bean evaporates and puffs out the shell, making it separate much more easily when the bean breaks during the cracking procedure.

In the case of nib roasting, the machines used are very similar to those for bean roasting. For liquor roasting, however, the nib must be finely milled to turn it into a liquid. This requires very careful moisture control. If the moisture content is too high, the cocoa liquor will be a thick paste and not a liquid. Even small amounts of water react strongly with this cellulose–protein–fat system, and a moisture content of about 10% would produce an almost solid material which would be very difficult to grind. Very low moisture contents would, on the other hand, produce a chocolate with a poor flavour. During roasting the flavour precursors can react in different ways, depending on the amount of moisture present. At very low moistures they are unable to produce the desirable reaction compounds.

### 3.2.5 Roasters

The process may be a batch or continuous one. Instead of the ball-shaped roaster (Figure 3.1) a drum shape is more commonly used (Figure 3.5). These can process as many as 3 tonnes of beans in a single batch. The heat can be applied externally through the walls or by passing hot air through the drum.

In order to kill the microbiological contaminants, both water and heat are required. These roasters are designed to add water or steam so as to increase the effectiveness of the kill. Care must be taken, however, to dry the beans again before the proper roasting takes place, as too much water can remove desirable as well as undesirable flavour components. As mentioned previously, too dry beans are also undesirable. Normally the temperature is raised to its roasting level of between 110 °C and 140 °C when the moisture level has been reduced below 3%. The total roasting procedure normally lasts between 45 minutes and 1 hour. After roasting, the product is usually cooled in an external cooler.

Where it is necessary to process large quantities of beans or nibs, a continuous process is often used. A typical continuous roasting...
system is illustrated in Figure 3.6. Here the beans are fed in batches through the top onto a shelf system. This shelf is made up of a series of slats, through which the hot air blows. After a predetermined time, the slats tilt in turn, starting at the bottom one, until the top one has returned to its position and a new batch has been fed into the top. In this way the beans fall onto the shelf below and through the roaster. The bottom shelves are used to give rapid cooling. Very large volumes of hot air pass through these roasters and care must be taken to ensure that this does not take with it some of the more volatile flavours, together with the moisture that evaporates during the roasting process.

Liquor roasting is carried out in specially designed equipment, which spreads the hot, liquid cocoa liquor over the surface of a long, hot cylinder with a fast rotating central column. Rotors and paddles, attached to the column, continuously stir the liquid and scrape the surface to stop the cocoa from becoming too hot. This process may take as little as 1 or 2 minutes.

### 3.2.6 Chemical Changes during Roasting

Unroasted beans usually taste very astringent and bitter. The high temperatures and drying during roasting remove many of the volatile acids, especially ethanoic acid, and make the nibs, or beans,
taste less acidic. The less volatile acids, such as ethanedioic (oxalic) and lactic, remain largely unchanged by the roasting process.

3.2.7 Maillard Reaction

This reaction, also called non-enzymic browning, is important to food quality throughout the food industry and gives products their colours and flavours when they are baked, toasted or roasted. It is an extremely complicated reaction involving many low molecular weight components with hundreds of different reactions and intermediate products. These intermediates may or may not have a

Figure 3.6  Diagram of continuous bean/nib roasting system (Lehmann Maschinenfabrik GmbH, Germany). (A) Product feed; (B) feed rollers; (C) exhaust air fan; (D) air heater; (E) air filter; (F) extraction screw. (Heemskerk1)
flavour of their own. Some act as catalysts for other reactions, whereas others may stop a particular series of reactions taking place.

Heat is required for the reaction to take place at a significant rate. Its products can be seen and tasted when food is burned onto a pan, the contents of which have not been stirred well enough. Water must also be present together with a reducing sugar, such as glucose, and an amino acid, peptide or protein. In the case of cocoa, about 12–15% of protein is present before fermentation. But the heat and acidic conditions during this treatment break down much of this into amino acids, thereby forming some of the precursors, which are converted by roasting into chocolate flavour.

The principal pathways of the Maillard reaction are shown in Figure 3.7. All the reactions take place at any pH $> 3$, but the actual pH does alter the probability of their taking place. The reactions on the left-hand side are largely flavour formation routes. The sugar splits up into smaller carbon chains. ($C_1$, $C_2$ etc. denote the length of the chains). The key intermediates 1DH, 3DH and 4DH are 1-, 3- and 4-deoxyhexosulososes respectively and are dicarbonyl compounds. The pathways on the right-hand side are often involved in colour rather than flavour production.

![Figure 3.7 Model of the Maillard reaction.](image)
The Strecker reaction involves the formation of aldehydes, some of which form part of the chocolate flavour, from the amino acids, which are largely tasteless and odourless. One of these reactions involves the reaction of an amino acid called glycine with glyoxal (a 1,2-dioxo compound). These eventually form pyrazines, which are heterocyclic compounds with two nitrogen atoms in a six-membered ring. The amounts of the different pyrazines formed depend very much on the temperature and time of the roasting reaction. Their measurement has, in fact, been used as a method of determining the degree of roast of cocoa liquor (see Chapter 8).

The characteristic smell of chocolate can also be produced by the reaction of amino acids such as leucine, threonine and glutamine with glucose, when heated to about 100°C. Higher temperatures will produce a much more penetrating/pungent smell.

3.3 GRINDING COCOA NIB

There are two aims when grinding the cocoa nib. The first is to make the cocoa particles small enough so that they can be made into chocolate. There is further grinding during the later chocolate making process, so it is not necessary to mill the nib very fine at this stage. The second more important reason is to remove as much fat as possible from the cells within the cotyledons. The fat is needed to help the chocolate flow, both when making the sweets and also when it melts in the mouth. The fat is also the most expensive major ingredient in chocolate, so economically it is necessary to make the most use of all the fat present. The fat is contained in cells which are on average between 20 and 30 microns long and from 5–10 microns wide/high.

Figure 3.8 shows sections through two beans as viewed under a microscope. The fat has been stained so that it appears dark. When moisture is present with fat, the two cannot mix, but they can exist together in two forms: i.e. a water-in-oil emulsion, with oil/fat surrounding water droplets; or an oil-in-water emulsion in which the oil/fat is in small droplets. The stability of these small droplets is aided by certain types of phosphoglyceride (also known as phospholipid), which form a surface between the water and fat. These are called emulsifiers (see also Chapter 5). Within the cocoa nib there is an emulsifier called lecithin. The two types of emulsion
can exist within the cells of cocoa. Figure 3.8a showing a largely oil-in-water emulsion and Figure 3.8b the reverse phase.

The aim of the milling is to get the fat from within the cells, so that it can coat the solid non-fat particles within the chocolate. Fat can be released by breaking the cells open. There is more fat within the cell than is necessary to coat any new surfaces created by tearing open the cells. This means that as the grinding reduces the cell size the cocoa liquor produced becomes thinner – there is more free fat around. Eventually there is no more fat to be released, and further milling creates only new surfaces to be coated with fat as it tears apart the cells. This makes the cocoa liquor become thick again. Some experimental figures that illustrate this are shown in Figure 3.9.

Figure 3.8  Sections through cocoa beans as seen through a microscope (fat is stained black). (a) Oil-in-water emulsion; (b) mainly water-in-oil emulsion.
The cell walls are largely composed of cellulose. The rate that fat can pass through the cellulose depends on the amount of moisture that is present (see Project 3 in Chapter 12). When fat is being pressed out of the cocoa for the production of cocoa powder, this extraction may be aided by the addition of some moisture to the liquor. When grinding, however, as was noted for liquor roasting, it is better to have a lower moisture to enable the cocoa liquor to flow more easily.

### 3.3.1 Cocoa Mills

It is necessary to grind the nib from a maximum particle size of about 0.5 cm down to less than 30 microns. This means that the particles must be ground down by 100 times. Most milling machines can only operate efficiently if the reduction is 10 times, so at least two grinding stages are required. In addition some mills work better with hard material, whereas others will work only with liquids. This means that cocoa is normally ground twice, initially by an impacting mill which melts the fat and produces a liquid containing large particles several hundred microns in diameter. The second mill is often a ball mill, which will work only with a liquid, or a disc mill, based on the original corn mills, which will work with liquid or solid material. Milled cocoa particles include cocoa starch, which make up about 7% by weight of the cocoa liquor. This has a particle size of between 2 and 12.5 microns and so is not

![Figure 3.9](image)

**Figure 3.9** Viscosity of cocoa liquor ground to different finenesses.
destroyed by the milling process. About 10% of liquor is made up of cellulose and a slightly larger percentage is protein.

If the factory is going to press the cocoa liquor to make cocoa powder, the cocoa liquor is not normally ground as finely as when the cocoa liquor is used for chocolate. This is because very finely ground cocoa liquor particles will block the filters in the cocoa presses and make it more difficult to remove the cocoa butter. In chocolate making, however, it is advantageous to have as much cocoa butter as possible released from the cells.

3.3.1.1 Impact Mills. Impact mills work by hitting the cocoa nibs with fast moving pins or hammers. Sometimes the particles are hit against sieves or screens. The cocoa butter melts due to the heat from the impact and from the mill itself and any free fat, together with the smaller particles, pass through the sieve. The larger particles remain on the inside until broken by the next series of pins or hammers.

3.3.1.2 Disc Mills. Disc mills often consist of three pairs of carborundum discs (see Figure 3.10). The cocoa liquor or nibs are fed into the centre of the top set of discs, where one disc is rotating but the other one is stationary. The discs are pressured together and the cocoa mass is forced through them to the outside by centrifugal force. The high shear destroys a lot of the particles.

Figure 3.10 Schematic diagram of a triple disc mill (Lehmann Maschinenfabrik GmbH, Germany).
releasing a lot of the fat. The cocoa liquor then runs down a chute to the centre of the centre set of discs and then finally onto the third set.

3.3.1.3 Ball Mills. The majority of the world’s cocoa is ground using ball mills, which can grind only liquids and so are normally preceded by impact mills. The mills contain large numbers of balls, which are made to bounce against each other, either by tumbling them by rotating the wall of their container or by a centrally rotating shaft with rods placed at intervals at right angles, as illustrated in Figure 3.11. The balls impact and rotate (Figure 3.12) and any particles caught between them are broken by the crushing or pulled apart due to the shearing of the rotating action. The smaller particles move more quickly in the fat as it is pushed away by the moving balls, but the larger ones are more likely to be milled as they move more slowly. With coarse cocoa liquor containing particles several hundred microns in diameter, the balls may be as large as 15 mm. Where it is desired to have a finer liquor, a series of ball mills may be used, each one containing smaller balls, down to as small as 2 mm. More small balls will fit in the same space, so the chance of a particle being caught between two of them greatly increases. The agitator speed is also increased with the smaller balls.

Figure 3.11 Schematic diagram of a stirred ball mill.
Sieves are used on the outlet to the mills to stop the balls damaging machinery during the next stages of chocolate making. The balls wear and are replaced at regular intervals. Magnets in the outlet pipes trap any pieces of metal that come out of the mill.

3.4 COCOA BUTTER AND COCOA POWDER PRODUCTION

3.4.1 Alkalising (Dutching)

Most of the cocoa liquor that is used to produce cocoa powder is alkalised, whereas very little liquor that is used to make chocolate is treated in this way. The alkalising process was developed in The Netherlands in the nineteenth century, which is why it is also known as the Dutching process. The reason for doing this was to make the powder less likely to agglomerate or sink to the bottom when it was added to milk or a water-based drink. The actual ability of the alkali to do this is not entirely certain, but it does affect both the colour and the flavour.

A solution of alkali, typically potassium carbonate, is normally added to the cocoa nib before roasting, although the cocoa liquor itself, or even the powder, can also be treated. Care must be taken not to add too much alkali. This is because the cocoa butter molecule is
composed of three acids attached to a glycerol backbone (Chapter 6). These acids may react with the alkali to produce soapy flavours. To overcome this, small amounts of ethanoic acid or tartaric acid may be added after alkalisation, in order to lower the pH. For certain types of bean which taste very acidic due to the presence of ethanoic and other acids in the cocoa, a mild alkalisation to neutralise these acids can be very beneficial to the eventual cocoa/chocolate flavour.

The reason for the change in colour is due to reactions based on a class of chemicals found in cocoa called tannins (polyhydroxyphenols). These are made up of epicatechin molecules (Figure 3.13), which during the different fermenting, drying and roasting stages may join together, oxidise or react with other chemicals within the cocoa. This increases the number of colour-giving molecules and makes the cocoa much darker. By carefully adjusting the pH, moisture, roasting temperatures and times, it is possible to produce a wide variety of colours.

### 3.4.2 Cocoa Butter

The highest quality cocoa butter is obtained by pressing cocoa liquor in a horizontal press of the type shown in Figure 3.14. The top section (4) consists of a series of pots, the base of each being a stainless steel mesh. Hot cocoa liquor is fed into the pot, which is then pressed by a steel ram operating at a pressure of about 40–50 MPa. The nib initially contains about 55% cocoa butter and this pressure is capable of pushing more than half of this fat through the sieve, where it flows down a pipe to the weighing point. The pressure is initially applied slowly, to prevent the formation of hard layers that would stop further fat from coming out. It is possible for the operator to
increase the pressure until the required amount of cocoa butter has been removed. This leaves a hard material in the pots, which contains between 8% and 24% of fat, depending on the type of powder being produced. The hard round discs of material are known as cocoa press cake. When the pots are opened, these are automatically ejected and fall onto a conveyor belt (9).

A lower quality butter is obtained by a continuous expeller process which presses the fat out of whole cocoa beans, which includes the shell. Very often these beans are not of an adequate standard to make chocolate, in that they may not be properly fermented or may have too acidic a flavour. The shell contains some fats which are not cocoa butter. These too are pressed out and get mixed with the cocoa butter. This is detrimental to the hardness and setting properties of the cocoa butter because of its eutectic effect (see Chapter 5). Very often this cocoa butter is cloudy and must be filtered.

Where the press cake is produced from whole or inferior beans, it has little value and is normally used as animal feed. Alternatively, the remaining fat can be removed by solvent or supercritical fluid extraction.

When buying cocoa butter, a series of specifications are given to the producer. These include a maximum free fatty acid content (usually 1.75%). This is where acids have come free from the glycerol “backbone” of the triglyceride. These will upset the setting properties of the chocolate. Also there is a maximum
saponification value (0.5%), to ensure that it does not give a soapy flavour. (The saponification value is the number of mg of potassium hydroxide (KOH) required to react with 1 g of the cocoa butter).

Pure pressed cocoa butter has a flavour, which will become part of the whole chocolate. For some products, in particular, white chocolate, this flavour is regarded as unpleasant. In this case deodorised cocoa butter is used. This is often produced by steam distilling the cocoa butter under vacuum.

3.4.3 Cocoa Powder

Cocoa powder is produced by milling the cocoa press cake. Once this leaves the presses it is broken into pieces less than 3 cm in diameter between two spiked rollers rotating in opposite directions. A cooled pin mill is then used to finely grind the powder. This must then be strongly cooled as it is transported in an air stream through a long pipe to the packing area. Most of the fat is still liquid after the mill and this must be solidified before packing in order to prevent the powder from sticking together. It is then collected in a cyclone separator with the finer particles being removed by a filter system.

Most powder is produced with a fat content of 20–22%. Lower fat ranges are available, e.g. 15–17% or 10–12%. A fat-free powder is also produced and sold for low fat or fat-free products.

The powder can be mixed with other fats to produce chocolate flavoured (compound vegetable fat) coatings or to produce cake mixes and fillings etc. A very large amount is used to make chocolate drinks. These are made from sugar, cocoa powder and lecithin. The lecithin can be added to the broken press cake and then milled with it. This makes sure that the lecithin is closely bound to the cocoa, especially the fat. Lecithin acts as an emulsifier and forms a boundary between the fat and other cocoa particles and the water when the drink is made. This helps these particles to disperse throughout the water rather than forming lumps.

REFERENCES


CHAPTER 4

Liquid Chocolate Making

Most people think of chocolate as a solid, because this is how they buy and eat it. To the chocolate maker, however, it is normally a liquid and is only solidified just before it is ready to be packed and sent to the warehouse or shop.

It has already been noted that dark chocolate is made mainly from sugar, cocoa nib and cocoa butter. The basic ingredients for a typical dark chocolate, roughly in the proportions that they are present, are shown in Figure 1.2. Similarly the basic recipe for a milk chocolate made from full cream milk powder is shown in Figure 4.1. As can be seen, these particles are relatively large, some being several millimetres in diameter. As with cocoa liquor, these must be milled so that the largest particles are smaller than 30 microns.

The fine particles must then be coated with fat so that they can flow past one another when the chocolate melts in the mouth. This process takes place in a machine called a conche (see Chapter 1). This machine is the only one specifically designed for the chocolate making process. The mixers are found throughout the food industry, whereas the mills are used to grind many food and non-food products (such as printing inks).

The liquid chocolate is then used to make the final product. This normally takes place by pouring it into a mould or passing it through a curtain of chocolate in a machine known as an enrober (Chapter 7).

How the chocolate flows during these processes is very important in order to produce the correct weight and appearance. The viscosity of the chocolate varies depending on how vigorously it is stirred or poured, and it is what is known as a non-Newtonian liquid (Chapter 5). Its behaviour is often described by two viscosity
parameters known as a yield value (related to the energy to start it flowing) and the plastic viscosity (its thickness when it is moving relatively quickly). Both of these parameters must be correct, otherwise waste product may be made. For example, “feet” may form on the base (Figure 4.2) and result in the product being sold at a low price as a misshape, or there may be holes, which will leave the centre exposed (Figure 4.3). Where the centre is moist it will dry out very rapidly, because it is no longer protected by the chocolate and its shelf life will be very much shorter.
The next chapter looks at the factors that control chocolate flow and which are used by the manufacturer to obtain the correct viscosity for subsequent processing.

### 4.1 CHOCOLATE MILLING

The aim of this process is to make sure that there are no particles in the chocolate that will make it taste gritty, in other words so that there are no particles larger than 30 microns. Another important factor is to ensure that there are not too many very small particles. Unlike cocoa liquor (see Chapter 3, Figure 3.9), which becomes thinner on grinding, chocolate becomes thicker. This is due to the creation of more fine particles, as will be explained in more detail in Chapter 5.

There are two different methods of grinding the chocolate ingredients: namely fine (or separate) ingredient and combined milling. In the fine ingredient process the solid, non-fat components are milled separately and then added to the cocoa liquor, cocoa butter and other liquid ingredients in the conche. In the combined milling, these ingredients are mixed with the cocoa liquor and some of the other fat before milling takes place. The two processes are likely to give a different flavour, as the sugar will pick up many of the aromas in the mill where it is being ground, and in the latter process there is cocoa in close proximity.

![Figure 4.3](image-url) Chocolate with part of centre uncoated.
Each process has its own advantages and disadvantages apart from the flavour aspects. The fine ingredient process is better able to control the number of fine particles, but the particles are largely fat free at the end of grinding. This means that the fat coating process in the conche takes longer than if many of them were already coated, as is the case for combined milling.

4.1.1 Separate Ingredient Grinding Mills

The mills are required to reduce sugar and other solid particles by a factor of approximately 100, from millimetres to tens of microns. This is a similar ratio to breaking a brick into pieces the size of sugar grains. When making cocoa liquor it was noted that it was best to reduce particle size in a series of steps, rather than in one operation. This is also true for chocolate ingredients. The hammer/pin mills of the type used to break the cocoa nib are very effective in shattering sugar. This is a very brittle material and when hit hard by a fast moving metal hammer or pin will break into a lot of smaller pieces. Milk powder is a lot more elastic and difficult to break and so requires much longer in the mill.

The traditional method of making chocolate was to pass the sugar through such a mill so that the mean particle size was about 100 microns. This was then added to the milk components and the cocoa liquor for combined grinding. This is still carried out in some factories, but the process has largely been superseded by a combined grinding process using two sets of roll refiners, as described in the next section.

In order to reduce the particle size still further to the required chocolate particle size, another milling stage is normally required. However, it is possible for the total process to take place in a single machine, called a classifier mill, one type of which is illustrated in Figure 4.4. Within it a series of milling stages can take place before the finely ground particles leave the mill.

The sugar and dried milk particles are fed into the mill via a chute (1) onto a milling disc (4). This rotates at several thousand times a minute and has metal hammers, wedges or pins at the edge (3), which hit the particles, breaking some and chipping pieces off others. A large volume of air is blown through the mill, entering at (6) and blowing out at (5). This lifts the particles and tries to pull them through the classifier (2). This consists of a rapidly rotating
A hollow cylinder with slits cut in the side. In fast flowing air, the smaller particles can travel at almost the same speed as the air, but the larger ones are much slower because of their weight and inertia. As the air passes through the slits, the smaller ones are able to pass with it and travel out of the mill, where they are collected by cyclone separators and filter bags. The larger ones, being slower, are hit by the moving bars between the slots and sent back into the breaking zone again. The larger particles will recirculate for as many times as is required for all of them to be small enough to pass through the classifier. In practice this means that any small sugar particles which are chipped or smashed off the larger crystals pass out of the mill at the first circuit, but some of the sugar and most of the milk will recirculate several times. All the particles within the mill travel at high speed and bump into each other, causing a lot of additional breakage.

There are two controls over the particle size that the manufacturer can use to obtain the one needed for the chocolate being
made. One is the rate of airflow. If this increases, more particles are pulled through the slits and a coarser product is made. If the speed of the classifying cylinder is increased, however, the reverse happens, the bars catch more particles and the product becomes finer. Because the particles pass from the mill as soon as they have been reduced to the required size (unlike combined milling, where all the particles pass through the mill until the largest ones have been destroyed), this type of mill produces relatively fewer finer particles than combined milling.

The mills generate a lot of heat, and this can cause some of the sugar to turn from a crystalline into an amorphous form (see Chapter 2). Also any fat present will melt causing the particles to become very sticky and block the pipes. When more than about 12% fat is present some form of cooling is required. At higher levels still, liquid nitrogen can be fed in with the air, and cryogenic grinding carried out. This type of mill is often used to process cocoa powder from press cake, as well as to manufacture chocolate.

4.1.2 Combined Milling

In the process used by many modern chocolate factories, the solid particles are milled using roll refiners. The process is shown schematically in Figure 4.5.

Initially the cocoa liquor, granulated sugar and milk components are placed in a heavy duty mixer together with some of the cocoa butter. (For chocolates made with a crumb, most of the solid

Figure 4.5  Diagram of two-roll and five-roll refiner chocolate making plant.
components are contained within the crumb, which is in the form of a partly milled powder.) It is important to turn this mixture into a uniform paste with the appropriate consistency in order for the milling to proceed correctly. This paste is then fed into a two-roll mill. This consists of two cylinders, placed horizontally side by side, which turn in opposite directions so as to pull the paste into the gap between them. If the paste has the wrong texture it will just form a bridge between the two rolls and the process stops. If it is correct, the pressure and shear in the gap will break some of the particles and also coat some of the newly formed surfaces with fat, so that a drier paste is formed, with a maximum particle size of between 100 and 150 microns.

The final grinding takes place on a five-roll refiner, similar to the one shown in Figure 4.6, which can be between 75 cm and 2.5 m wide. These take the paste down to a maximum particle size between about 15 microns and 35 microns. The actual size will depend on the type of chocolate being made, and greatly affects its flow properties as a liquid, as well as the taste and texture in the mouth (see Chapter 5).

The five-roll refiner consists of five slightly barrel-shaped horizontal cylinders, with four of the cylinders placed one above the other (Figure 4.7). The first, or feed cylinder is placed below the
others, but on the side so that a trough is formed between it and the second cylinder, which will contain the paste from the two-roll refiner. As it has four crushing gaps and is grinding so finely, the five-roll machine operates much slower than the two-roll one, with its single gap. Because of this, a two-roll machine is normally used in conjunction with several five-roll ones.

The cylinders are hollow and can be cooled or heated by water which flows through them. They are also pressed closely together, usually by a hydraulic system. This pressure bends the barrel shape so that it becomes straight and there is a uniform straight gap between the cylinders. A knife blade placed against the back of the fifth cylinder removes the chocolate in the form of flakes or a powder.

The big particles can be broken in many ways. Hitting them hard, as in the hammer mill, can break them into two or more similar sized pieces, or can chip smaller pieces off the edges. Crushing between two hard surfaces was shown to be one of the ways ball mills operate when milling cocoa liquor (Figure 3.12). Yet another way is to use shear to pull them apart. Shear is related to the difference in speed between two moving surfaces divided by the distance between them. This means that if two surfaces are travelling at very different speeds and are very close together, there

Figure 4.7  Schematic diagram of a five-roll refiner.
is a very high shearing action, which will pull the particles apart. This is in fact the case with a five-roll refiner.

Each successive roller is faster than the previous one (Figure 4.7), and because the film of chocolate is attracted to the faster moving surface rather than the slower one, it continues going up the refiner rather than keeping going round and round the bottom one.

This type of machine works by having a continuous film of chocolate from the feed trough (hopper) to the knife. The thickness of the film depends on the gap between that particular roller and the one below it. What actually takes place is illustrated for the gap between the second and third rollers in Figure 4.8. The lower roller is turning at say 55 rpm and has a film thickness of 100 microns. The second roller is turning at 150 rpm, but the film has to remain continuous. This means that it is stretched out by the higher speed so the thickness is reduced according to the relationship between the speeds of the two rollers, i.e. it becomes $\frac{100 \times 55}{150}$ microns thick ($=37$ microns). The final fineness therefore depends on the ratio of the speeds of the different cylinders together with the thickness of the initial film. This latter depends on the outlet width of the slit from the trough, i.e. the gap between the first two rollers. Strangely it is affected very little by the pressure, which mainly acts to give a uniform film along the roller.

The temperature also plays an important part in the operation of a roll refiner. This alters the texture/viscosity of the film by changing the flow properties of the fat present. Because the roller

![Figure 4.8](image) Close-up of the gap between the second and third rolls of a five-roll refiner.
surfaces are turning at relatively high speeds, there is a centrifugal force on the individual particles trying to throw them away from the machine. The film itself is pulling them on, as long as they remain part of the film. If something goes wrong with the texture of the film, e.g. some of the fat sets because it is too cold, the particles become free and the chocolate is thrown away from the machine. Temperature control is therefore very important when grinding chocolate.

The shear between the rollers not only breaks the particles, it also coats some of the newly created surfaces with fat. In addition, as the breakage occurs, the newly formed surfaces, which are chemically very reactive, are able to pick up the volatile flavour chemicals from the cocoa particles being broken nearby at the same time. This means the chocolate is likely to have a different flavour to one made using the separate grinding process.

4.2 CHOCOLATE CONCHING

The chocolate conche was invented by Rudi Lindt in Switzerland in 1878 and was named after the shell, which it resembled in shape. He said that it helped make his chocolate smoother and that it modified the taste. When the conche was invented, the ability to mill chocolate was poor, so it is possible that particles were broken in his conche, which made the chocolate smoother. Today, however, the grinding systems are very efficient and almost no further breakage takes place, other than when particles are loosely held together as agglomerates. The conche still does change the flavour of the chocolate and also the way it melts in the mouth. Additionally, of critical importance to the chocolate manufacturer, it determines the final viscosity of the liquid chocolate before it is used to make the final products.

The conching process is in fact two distinct processes which take place within the same machine. The first is flavour development. The fermentation and roasting processes produce the flavour components required to give chocolate its pleasant taste, but they also result in some undesirable astringent/acidic ones that it is necessary to remove. In addition, some chocolates need further flavour development, for example, for some purposes, an enhanced cooked flavour is desirable.

The second process is to turn the chocolate from a powder, flaky or thick dry paste into a free flowing liquid that can be used to
make the final products. This involves coating the surfaces of the solid particles with fat, so that they can slide past one another.

4.2.1 Chemical Changes

During fermentation, ethanoic acid and to a lesser extent other short-chain volatile fatty acids such as propionic and isobutyric (2 methyl propanoic) acids are formed. However they have boiling points above 118°C, which is considerably higher than the temperature of the chocolate during most conching procedures. Moisture is, however, removed from the chocolate, particularly during the early parts of the process (Figure 4.9), and this may aid the removal of the acids by a type of steam distillation process.

Other workers have noted a large reduction in the amount of phenols during the first few hours of conching. It is not known, however, what if any effect these compounds have on chocolate flavour. Headspace analysis of the air above a conche has shown that the amount of volatiles decreases by 80% during the first few hours of conching. It is also possible to over-conche. Chocolate that has been processed for too long may have very little flavour at all.

The chocolate flavour depends on the time and temperature used, in general a higher temperature means a shorter processing time. Above about 70°C in milk chocolate however, cooked flavour changes start occurring. Some manufacturers use temperatures above 100°C to try to promote some Maillard type flavours. Because there is very little water present, these flavours are not as

![Graph showing the changes in moisture and acidity during a conche cycle (time in hours). (Beckett)](image)
strong as those developed during high temperature milk drying or the crumb making process. In some milky flavoured chocolates these Maillard flavours must be avoided so the conching temperatures must be kept below 50 °C. This is also true for some sugar-free chocolates containing sugar alcohols. Here higher temperatures melt the crystals and they later resolidify in gritty agglomerates.

4.2.2 Physical Changes

Recent research by Dr Ziegleder\textsuperscript{2} in Germany has shown that a major process that takes place in a conche is the physical movement of the flavour molecules between the different components of the chocolate. At the start of conching the chocolate flavours are only on cocoa particles and in the cocoa butter, and the milled sugar has only sweet flavours (see Figure 4.10). During conching, cocoa flavour and fat are partially transferred onto the sugar surface. This coating results in a more uniform cocoa flavour and less sweetness than is found with freshly roll-refined sugar.

![Figure 4.10](image_url)

Flavour distribution between cocoa solids, sugar particle surfaces and the fat phase before and after conching. (Ziegleder\textsuperscript{2})
movement of flavour is caused by the initial flavour concentration gradient between the different particles or phases. In addition any amorphous sugar formed during roll refining (Chapter 3) may help absorb the flavour.

The transfer was examined under different conching procedures and then the chocolates produced were examined by chemical and sensory analysis. In general a preference was found when this coating action was optimised by carrying out more of the conching when the chocolate was thick and pasty, rather than when it was thin and runny.

4.2.3 Viscosity Reduction

This process is essentially one of coating the particles with fat. As with grinding, shear is an important factor. Figure 4.11 illustrates some particles at rest and then under shear.

The shear in this case is once again related to the difference in velocity of the surrounding walls divided by the distance between them and we can define a shear rate as:

\[
\text{shear rate} = \frac{(v_1 + v_2)}{h}
\]  (4.1)

Once again high speeds or narrow gaps have a bigger effect on the particles between them. What we are trying to do is to put fat over

![Figure 4.11](image1)

**Figure 4.11** Representation of simple shear between two flat parallel surfaces. (Beckett³)
the surface, which is a bit like trying to butter bread, where the fat between the knife and the bread is subjected to a high shear and so is forced into a thin layer over the surface.

In chocolate making this coating action is particularly important because it is desirable to make the chocolate flow as well as possible (have the lowest possible viscosity) at any particular fat content. Figure 4.12 illustrates the same chocolate (identical fat content) processed at different shear rates. As can be seen, all reach different equilibrium viscosity readings and they will retain these for however long they are sheared. This means that within the normal conching shear rates used, the higher the shear the thinner the chocolate will be. This however requires very large motors and a lot of energy and there is a practical limit to the shear that can be developed.

There are two approaches to reducing the chocolate viscosity. One is to have a very large stirred tank, where only a small amount of the chocolate is being sheared at any one moment. Because there is a lot of chocolate in the tank, it is possible for the chocolate to be inside the tank for several hours and for there still to be a throughput in the region of tonnes per hour. The other approach is to highly shear a few kilograms at a time in a continuous processor. Because of the small amount inside the machine, it can stay there for only a fraction of a minute to give the same throughput as a large conche.

Another thing that this process does is break up groups of particles that are loosely stuck together. These agglomerates can take two forms. In one of them (Figure 4.13a) there is no fat within it, so the breakage just gives new surfaces that have to be coated with fat and so the viscosity is increased. In the other (Fig 4.13b)

![Figure 4.12](image)

**Figure 4.12** Change of viscosity with time for conches with different shearing actions. (Beckett³)
there is fat in the middle that is surrounded by the solid particles and so does not aid the flowability of the chocolate. Once this is broken, however, more fat is released than is necessary to coat the new surfaces, so the overall viscosity is reduced.

4.2.4 Conching Machines

4.2.4.1 The Long Conche. This was the type of machine that was developed by Rudi Lindt and consisted of a granite trough containing a granite roller (see 1.5 and Figure 4.14). The roller pushed the chocolate backwards and forwards for a long period of time, often amounting to several days. As the surface was changed, the volatile flavour components were able to escape into the air. Very often these were loaded from large wheelable tanks using a shovel. This was very hard and hot work and in the early part of the twentieth century the life expectancy of people working in that part of the factory was relatively short. Usually four troughs were attached and operated together from a single motor, or from a pulley which was driven by leather belts from a central shaft which ran the length of the conche rooms. Much of the early machinery was belt driven and some factories had saddlers’ shops to maintain the leather.

The density of the powder that is fed into the conche is less than half that of the finished chocolate, so the troughs appear relatively empty at the end of the process, even though they may have been nearly overflowing at the beginning.
These conches have poor temperature control and a high energy consumption together with a relatively small capacity and so have almost entirely been replaced by more modern designs.

4.2.4.2 Rotary Conches. This type of conche is so named because the mixing elements rotate within the tank body, which forms the outside of the conche. Many of the earlier designs (Figure 4.15) were round with a vertical central shaft which drove the mixing or scraping arms. Others had additional mixing elements which performed a planetary motion around the centrally driven arms. Most were open topped to enable the moisture and flavour volatiles to escape. As with the long conche, they tended to have poor temperature control and were usually limited in size to about 1 tonne capacity, although 3 and 5 tonne machines were made. They have therefore been largely superseded by conches with horizontal stirring elements.

A typical modern conche is illustrated in Figure 4.16. Here the tank has three connecting troughs and three stirring arms. As they rotate, they smear the chocolate against the side of a temperature controlled wall – the “buttering bread” action. The machine then throws it in the air, which enables the volatiles and moisture to escape more easily. The previous forms of conches normally could
not do this as they tended to compact the chocolate into the bottom of the conche.

The ends of the conche arms are wedge shaped. This means that when the chocolate is a thick paste, the arm can operate with the point first. The point will be able to cut into the paste and the side of the wedge can smear the chocolate against the wall. When the chocolate becomes thin, on the other hand, it will just flow around the wedge and not much mixing or coating of the particles takes place. By reversing the direction of the arms at this stage however, the flat end of the wedge causes much more movement and mixing to take place.

Many of these conches have louvres on top (Figure 4.17), which can be fitted with fans to aid volatile removal, if required. This makes them safer and more hygienic. Conches often process between 5 and 10 tonnes of chocolate in a period of less than 12 hours. The conches are filled automatically from conveyor belts feeding from the milling stage. They also empty automatically through pipes in the base. This is just one more illustration of how in about 20 years the chocolate industry has changed from being a highly labour intensive almost craft industry, to a modern, high throughput, large machinery one.
4.2.4.3 Continuous Low Volume Machines. This type of machine tends to be used to produce a lower viscosity chocolate once the flavour changes have taken place (although twin screw extruders have been used to carry out both processes). A typical liquefying machine is illustrated in Figure 4.18. It contains a central shaft with pins attached to it at intervals and which rotates at a high speed. The walls have pins attached which do not move (stators). The gaps between the pins are small and the velocity across this gap is very high, i.e. there is a very high shear rate. As the chocolate is
pumped along the pipe, it is subjected to this violent action, which causes the fat to coat much of the solid surfaces. The energy input is very high however, which makes the chocolate become hot. This would alter the flavour of the chocolate were it not for the pipe walls, which are cooled by cold water.

This type of machine can be used after conching or in parallel with it, i.e. by having some of the chocolate pumped out, liquefied
and then returned to the conche. In both cases the aim is to produce a thinner chocolate in a shorter time.

4.2.5 The Three Stages of Conching

To get a well processed chocolate it is normally desirable to let it pass through three stages:

1. dry conching
2. pasty phase
3. liquid conching.

In the dry phase the chocolate is still powdery and, for milk chocolate in particular, contains an excess of moisture. This is detrimental to the chocolate’s flow properties (see Chapter 5). In addition when it is removed it is able to take with it some of the undesirable acidic flavours (Figure 4.9). When a lot of the surfaces are still uncoated with fat, it is a lot easier for the moisture to escape. This means that if the chocolate is heated and mixed in the powdery state, the whole process can take place more quickly and a lower moisture, thinner chocolate produced.

As the temperature rises, more of the cocoa butter melts and the particles begin to stick together. Sometimes they form into balls several centimetres in diameter, which run around the conche before joining together to form a thick paste. Within the paste there are still a lot of milk and/or sugar particles that are still not coated with fat. When the paste is thick there is a relatively high probability of the shear/smearing action coating them with any fat that is nearby. Once it becomes thin however, these uncoated particles will just flow out of the way. In order to make a chocolate which flows well in the mouth (when the fat melts due to the body heat), it is necessary to coat as many of these surfaces as possible. This means that the paste should be kept as thick as it is possible for the conche motor to mix, for as long as possible.

The final function of the conche is to ensure that the chocolate has the correct flow properties for the next processing stages. This in turn will depend on the type of coating or moulding machinery being used. The final stage is therefore one in which the final additions of fat and emulsifier (see Chapter 5) are made to the
chocolate. This makes it very thin and little further mixing takes place.

The chocolate can then be pumped into storage tanks ready for use. Sometimes it is transported as a liquid in a road tanker to another factory. Alternatively the chocolate can be solidified and stored or transported as blocks or small chips.

REFERENCES

CHAPTER 5

Controlling the Flow Properties of Liquid Chocolate

The flow properties of liquid chocolate are important to the consumer and the confectionery manufacturer. Although there are many very sophisticated instruments for measuring viscosity and texture, the human mouth is really far more sensitive than most of these. When someone eats chocolate, the teeth bite through the solid chocolate. This means that the hardness of the solid chocolate is very important. The temperature of the mouth, at about 37°C, is above the melting point of the fat within the chocolate, so it rapidly melts, especially as it is subjected to the intense mixing and shearing of the teeth and tongue. Once it has melted there are two important factors. One is the maximum particle size. As was noted earlier, if there is a significant number of particles larger than 30 microns (0.03 mm) the chocolate will feel gritty on the tongue. In addition, for sizes below 30 micron a difference of 2–3 microns in maximum particle size can be detected as different levels of smoothness. Chocolates with a maximum particle size of about 20 microns have been sold as having a silky texture. The second factor is the viscosity. This not only affects the way the chocolate runs around the mouth, i.e. the texture, but it also changes the taste. This is because the mouth contains three different flavour receptors in different places (see Figure 5.1). The time the solid particles in the chocolate take to reach the receptors depends on the viscosity. This means that two chocolates made from identical ingredients, but processed to give different viscosities, will taste very different. (See Project 15 in Chapter 12). Particle size affects viscosity as well as texture, and a milk chocolate which
has been milled to say a maximum particle size of 20 microns will have a creamier taste and texture than one of 30 microns.

For the manufacturer, the weight control of the products is very important. Chocolate is a relatively expensive food in terms of both ingredients and processing. It is therefore economically important not to put too much chocolate on a confectionery centre. As will be shown in Chapter 7, the coating methods rely on the liquid chocolate having the correct viscosity. In addition, if the coating is not put on correctly, misshapes will be formed (see Figures 4.2 and 4.3), or the centre will be exposed reducing the shelf life of the product.

5.1 VISCOSITY

Viscosity has been described as “the resistance to motion when stirred or poured”. This is however, not necessarily described by a single number. Non-drip paint or tomato ketchup are everyday examples of materials with complex flow properties. The paint tin or the ketchup bottle can be carefully turned upside down with their lid removed and the contents will initially remain in place. If, on the other hand, they are vigorously stirred or shaken just before being turned, they will pour out as quite thin liquids. So how is viscosity defined?

We can consider viscosity or consistency as an internal friction to movement. When movement is easy (i.e. runny materials) there is
little friction, but for thick materials friction is high. For this it is useful to reconsider the idea of shear (see Figure 5.2). If the liquid has two flat surfaces each of area \( A \) and a distance \( h \) apart moving at velocities \( v_1 \) and \( v_2 \), then, from the previous definition, we have a shear rate (\( D \)) of \( (v_1 + v_2)/h \) (eqn 4.1). Because velocity has the units of length divided by time and we are dividing this by a length, \( h \), the shear rate must have the units of \( 1/\text{time} \) and is normally measured in \( \text{s}^{-1} \) (reciprocal seconds).

The force required to move the top plane relative to the bottom one is called the shear stress (\( \tau \)). If we plot the shear rate against the shear stress, in other words how fast the liquid moves as different forces push it, different curves are obtained, depending on which type of material was being measured (see Figure 5.3).

Viscosity (\( \eta \)) is defined as the ratio of shear stress to shear rate, \( i.e. \)

\[
\eta = \frac{\tau}{D}
\]  

(5.1)

The units of viscosity are Pascal seconds (Pa.s), although the older units of Poise (0.1 Pa.s) are also still used.

This is illustrated in Figure 5.3 where the viscosity is the gradient of the line, which for the liquid shown in curve 1 is the same for all shear rates. So if we double the force on this liquid it will move twice as fast. This is known as a Newtonian liquid, as it was first described mathematically by Sir Isaac Newton.

Substances such as golden syrup are Newtonian, but the vast majority of foods are non-Newtonian and follow a wide variety of other curves. Curve 2 shows a substance which does not move when smaller forces are applied, just as was the case when the paint
or ketchup was inverted. Once it starts to move, however, it behaves just like a Newtonian liquid. These substances are known as Bingham fluids.

Chocolate has a more complex flow behaviour, however. As with the Bingham liquid it takes a measurable force to start it flowing, but once it does so, the higher the force the thinner it becomes. This is a major problem for the chocolate manufacturer, because the viscosity is not a single number, but has a value which depends on how fast it is flowing. It can in fact be represented best by a flow curve.

In a factory it is not possible to use a curve as a specification for a chocolate viscosity, so this data has to be simplified. The way that this is normally done is to describe the curve by a mathematical equation. Many exist, but the most widely used is the Casson equation, which was originally developed to describe the flow of printing ink. The viscometer (see Chapter 8) measures a few points on the curve and then the equation is used to give two flow parameters: the yield value and the plastic viscosity.

The yield value is related to the energy required to start the chocolate moving. If it is high the chocolate will tend to stand up, which may be required for putting markings on sweets or when producing chocolate morsels for baked cookies. A low yield value is needed to give a thin coating of chocolate over a biscuit.
The plastic viscosity relates to the energy required to keep the chocolate moving once it has started to flow. This is also important in determining the coating thickness of chocolate on a sweet and also in determining the size of pumps needed to pump the liquid chocolate.

The remainder of the chapter describes how the chocolate processing and the ingredients can be used to adjust the viscosity of the chocolate.

5.2 PARTICLE SIZE

5.2.1 Particle Size Distribution Data

In the first part of this chapter it was shown that the viscosity of chocolate could not be described by a single number, but a minimum of two numbers was needed and, to be really accurate, a full curve was required. Particle size is the same. Although, so far, the maximum particle size has been referred to as determining the texture of a chocolate, very few particles are as large as this. Once again the real situation is a curve, known as a particle size distribution and the chocolate manufacturer has to take a summary of this information to aid with processing and quality control.

The particle size distribution can take several forms, two of the most common are illustrated in Figure 5.4. This shows the size distribution.

![Figure 5.4](image)

**Figure 5.4** Comparison of mass and number particle-size distributions of a chocolate.
distribution of the same chocolate but displayed in two ways. If the solid particles in the chocolate were viewed down a microscope and the diameter of each determined, it would then be possible to determine the proportion of particles within a given size range. This is shown in the number versus size distribution. If, on the other hand, the individual particles were weighed and the proportion of the total weight that was due to a certain size range was measured, then a distribution like the other one in Figure 5.4 would be obtained. This is a mass versus size distribution. Normally these are measured by determining an estimate of the volume of the solid particles and then assuming a constant density. (See Chapter 8 for details of the type of instrumentation capable of making this measurement.)

As can be seen, the two curves look very different. The mean particle size on the number distribution is about 2 microns, whereas it is almost 10 microns for the mass distribution. The maximum size on the number distribution appears to be about 9 microns, although it is known from the mass distribution that particles as large as 70 microns are present. Why is this so?

The difference is due to the fact that the number distribution is proportional to the particle’s diameter or radius \((r)\), whereas the mass relates to its volume, \(i.e.\) to the cube of the radius \((\text{volume} = \frac{4}{3}\pi r^3)\). A ten micron particle is therefore equivalent in mass to \(10 \times 10 \times 10 = 1000\) one micron ones on the mass scale, but the two are both equal to a single particle on the number scale. Although the large ones are present in the sample, on the number graph their proportion is too low to be significant and to be registered.

In chocolate a small number of larger particles will give a gritty texture, so the number distribution will not give the information that is needed. The mass distribution is better, but it is still not obvious what measurement should be taken to be representative of the large particles, \(i.e.\) the grittiness of the chocolate. Once again it is a distribution. Particles much larger than 50 microns may be present in the occasional sample, but it would be impractical to sample a lot of the chocolate. In fact the maximum particle size is going to vary very much from sample to sample within a bar of chocolate. The solution that is often used is to record the 90th percentile. That is, the size at which 90% of the solid particle mass is due to particles with a diameter less than this size. This appears
to correlate fairly well with what people actually taste and with measurements made of the largest particles using a micrometer (Project 6 in Chapter 12).

5.2.2 Effect of Particle Size on Viscosity

The largest particles are important for mouth feel with respect to grittiness, but the smaller ones are more important with respect to chocolate flow properties, in particular the yield value.

The reason for this is because a large amount of fat is required to coat them so that they can move past one another in the liquid chocolate. Figure 5.5 represents a cube of sugar. This has six sides (four vertical sides, the top and the bottom), which need coating with fat. If the cube is broken in half, then there are eight sides to be coated (the two new surfaces are the same size as the old ones). This means that there is a need for 30% more fat, although there is only the same amount of sugar present. As was noted in the

Figure 5.5  Sugar cube before and after being broken in half.
previous chapter, it is necessary to reduce the particle diameter by about 100 times, so a vast new surface is created, which will use up the fat which helps the chocolate to flow.

It is known that we need to mill the sugar below 30 microns, otherwise it would be rough. What would happen if it were possible to create spherical sugar particles all of which were 29 microns? This would have a minimum surface area to be coated with fat, but would have all the particles small enough to make chocolate. This, however, would give a very thick chocolate. The reason is that the particles would not pack together. As shown in Figure 5.6, all-spherical particles can fill only about 66% of the volume. If we take another size of specially chosen smaller ones to fill in the holes, this increases to 86%, and a third size will take this to 95%. So the chocolate manufacturer ideally would like to grind the particles so that they pack well together but have a minimum area to coat. At the moment it is only possible to make minor changes to the particle size by adjusting the operating speeds and conditions of the mills being used.

So what parameters have to be measured to determine whether the particle size distribution is a good one for chocolate making? The shape of the particle size distribution curves gives some indication, but, as with viscosity measurements, curves are difficult to use to control a manufacturing process. There is, however, one parameter which gives a useful guide as to how the particle size distribution will affect the flow.

![Figure 5.6](image)

**Figure 5.6** Packing of spheres that are: (1) all the same size; (2) two different sizes; (3) three different sizes.
Figure 5.7a represents a lump of sugar wrapped up as a present. If it is unwrapped and the minimum amount of paper has been used, we have a measurement of the surface area of the sugar (see Figure 5.7b). If all the “pieces of paper” from the particles within a defined volume are fitted together, an area per unit volume is produced, also known as a specific surface area. Machines are available to produce this measurement, which is normally expressed in m² cm⁻³. This measurement relates to the radius squared (the surface area of a sphere is πr²). So this measurement in fact is not obtained directly from either the number distribution or the mass distribution. A surface area distribution would lie between the two.

It has been shown that the fine particles make the chocolate thicker by using for coating some of the fat present in the chocolate that would otherwise enable the particles to flow past one another. They do not, however, affect both of the flow parameters in the same way. Figure 5.8 shows graphs of the yield value and of the plastic viscosity of the same chocolate that has been milled to different particle sizes. The yield value increases dramatically as the chocolate becomes finer, but the plastic viscosity stays almost unchanged, in fact decreasing slightly at one stage.

This is because over half the volume of the chocolate is taken up by solid particles, as illustrated in Figure 5.9.

When the particles are large there are a limited number of points of contact. As the number of particles increases, there are many points where the particles are near one another, so a loose structure is built up. This structure has to be broken before the chocolate will flow and so the yield value increases with the proportion of fine
Figure 5.8  Influence of particle fineness on the viscosity parameters of two milk chocolates: (1) 30% fat; (2) 32% fat. (Chevalley²)

Figure 5.9  Schematic representation of the solid particles and fat within a milk chocolate.
particles. Once the chocolate begins to flow, the structure is broken and the small particles can move along together. There is therefore very little difference between that and big particles moving, so the plastic viscosity remains almost unchanged. The slight decrease is probably due to more fat being released into the system by further grinding of the cocoa liquor or milk powder. As was shown in Chapter 3, cocoa liquor becomes thinner as it becomes finer, in contrast to chocolate.

### 5.3 EFFECT OF FAT ADDITIONS ON VISCOSITY

As would be expected, the addition of more liquid fat helps a chocolate to flow more easily (see Project 5 in Chapter 12). Milk fat has the same effect as cocoa butter on viscosity if added to chocolate at 40°C, but slows down the setting rate and softens the final chocolate (see Chapter 6). In addition, because milk fat melts at a lower temperature it will change how the product melts in the mouth. The two fats must therefore be present in the right ratio to give the correct chocolate texture in the product in which it is being used. Milk fat is often used in dark chocolate to delay the formation of a white sheen on the surface called chocolate bloom (Chapter 6).

The fat must be in a free form to aid the flow. The cocoa liquor and milk powder must be milled finely to release the fat from the cells or the spherical casing respectively. In addition the conching must have been sufficiently vigorous to break any fat-containing agglomerates.

Most chocolates contain between 25% and 35% fat, although ice-cream coatings are much higher (see Chapter 10), and some special products like cooking chocolate and vermicelli pieces are lower. The actual level present will depend on the process being used – a certain amount is needed so that the chocolate film remains on the roll refiner and so that the conche motor is not overloaded. In addition it will affect the texture of the finished chocolate, so a high quality tablet of chocolate is likely to have a higher fat content and a lower particle size than a chocolate that is used to coat a biscuit.

The effect of an extra 1% fat on the viscosity depends on the amount that is already there and also which of the viscosity parameters we are considering (see Figure 5.10). Above a fat
content of 32% there is very little change in viscosity with any further additions. A 1% increase to a 28% fat content has a dramatic effect especially on the plastic viscosity, which is almost halved. The change becomes more dramatic at even lower fat contents, as chocolates below 23% fat are normally a paste rather than a liquid, but 25% fat chocolates are available on the market.

The effect of fat is proportionately much higher for the plastic viscosity than the yield value (more than 12 times compared with less than 3 times for the samples and ranges illustrated). This is not too surprising as the extra fat will add to the free-moving fat that aids particles when they flow past each other. The majority of the fat is “wetting” fat, which is partially tied to the particle surfaces. This free fat has a large effect on lubricating the flow when it takes place and so the plastic viscosity decreases dramatically. The yield value is more connected with the force between the solid particles, which in turn is connected with the absolute distance between them. This will be less affected by the fat additions.
5.4 MOISTURE AND CHOCOLATE FLOW

It would be expected that as water is a liquid, if it were added to liquid chocolate the resultant mixture would have a viscosity somewhere between the water and the chocolate. This is far from the case and the addition of 3% or 4% by weight of water will turn chocolate into a very thick paste (Project 8 in Chapter 12). Very approximately, for every 0.3% extra moisture that is left within the chocolate at the end of conching, the manufacturer must add an extra 1% fat. Because the fat is by far the most expensive major component within the chocolate, it is important that as much “free” water is removed as possible.

Water is like the fat in that it can be bound or free; unlike the fat it should be as much as possible in the bound condition. If the total moisture is measured, for instance by Karl Fischer titration (see Chapter 8), then part of this is due to water of crystallisation in the lactose as it is often present as a monohydrate (Chapter 2). Other water may be inside the cocoa cells which have not been destroyed by milling. This water will not affect the chocolate flow.

If a bowl of icing sugar is left in a moist room it will soon form a lump due to the water sticking the particles together. Chocolate is almost half made up of very tiny sugar particles and any moisture will either dissolve them or form sticky patches on their surfaces. This causes them to stick together and greatly increases the viscosity of the chocolate. If large amounts of water are present, say above 20%, there is enough water present for it to dissolve most of the sugar and form continuous streams through the chocolate and so help it flow.

Cream can be added to chocolate to form a soft mixture, known as ganache, that does not have any snap when broken and does not contract very much on cooling. This is made by stirring the cream very vigorously as liquid chocolate is added to it. In this case the water in the cream is in an emulsion. This means that the water droplets are surrounded by a fat, with an emulsifier forming a layer between the two. These emulsifiers occur naturally in cream, but are also added to chocolate. When one of these emulsifiers is present, the chocolate viscosity is slightly less affected by the moisture.

Most of the water is removed from the conche during the initial dry conching stage (see Chapter 4). This must be done with care
however, because if the water vapour comes out of the chocolate ingredients faster than it escapes from the conche, it can condense to form droplets. These droplets can return into the chocolate and dissolve some of the sugar particles. These will stick together to form hard gritty lumps, once the moisture is eventually removed. This means that even if a chocolate has been correctly milled, it will still taste very sandy in the final product.

5.5 EMULSIFIERS AND CHOCOLATE VISCOSITY

The role of an emulsifier is to form a barrier between two non-mixable substances. An emulsion plays an important role in separating water globules in fat; for instance in margarine, which has water droplets in fat (a water-in-oil emulsion), or in cream, which is fat droplets in water (an oil-in-water emulsion). As was shown in Chapter 2, the fat within cocoa beans can exist in both types of emulsion. In chocolate there is almost no water, so the emulsifier is somewhat different. There are sugar particles, which are hydrophilic but lyophobic, in other words, which attract water but tend to repel fat.

Liquid chocolate flows because the sugar and other solid particles are able to move past one another, so, as was described for the conching process, the solid particle surfaces have to be coated with fat. This is something that does not occur very naturally, so, like the water emulsions, a substance that forms a layer between the solid particles and fat will greatly help the process. In this case, the emulsifier coats the solid surface and forms a boundary layer between it and the fat and is really a surface active agent rather than an emulsifier.

The mechanism by which a surface active agent works is illustrated in Figure 5.11. The individual molecules have a lipophilic (fat-liking) tail, which sticks out into the fat, where it wants to be. The other end is lipophobic and does not like the fat. It therefore keeps itself away from the fat as far as possible by attaching itself to the (also lyophobic) surface of the sugar. This can be compared with seaweed on a rock. There is a “head” which attaches itself to a rock, and a long “tail” which goes out into the sea. The “tail” waves around and alters the flow around the rock. Just as there are many types of seaweed, there are also many types of surface active
agent. Some have very large “heads” which bind very strongly to the sugar, whereas others are less strongly attached and may even be removed by the addition of a different surface active agent. Similarly there are different lengths of “tails”, which affect the flow properties in different ways. This means that a surface active agent that is especially beneficial with regard to the yield value may be poor with respect to the plastic viscosity, and vice versa.

5.5.1 Lecithin

The most common surface active agent is lecithin, which has been used in chocolate since the 1930s. It is a naturally occurring substance, frequently obtained from soya and regarded by many as being beneficial to health. As described above, it is able to attach itself to the sugar, whilst leaving the other end of the molecule free in the fat system to aid the flow. Harris$^5$ showed that the lecithin was able to bind itself particularly strongly to the sugar, and it is this phenomenon which makes it so effective in chocolate manufacture. This was later confirmed by Vernier$^3$ (Figure 5.12) who used a confocal laser scanning microscopy, which showed the fluorescing lecithin molecules to surround the sugar particles.

Additions of between 0.1% and 0.3% soya lecithin are said to reduce the viscosity by more than 10 times their own weight of cocoa butter. Also chocolates containing surface active agents,
such as lecithin, can tolerate higher levels of moisture than emulsiifier free ones. This is important because water is so very detrimental to chocolate viscosity.

Too much lecithin, however, can be detrimental to flow properties in that at higher levels, e.g. above 0.5% (Figure 5.13), the yield value increases with increased lecithin additions, although usually the plastic viscosity continues to fall. Bartusch\textsuperscript{4} showed that at 0.5% about 85% of the sugar was already coated. After this the lecithin may be free to attach itself to itself to form micelles, or form a bilayer around the sugar (so that the “tails” of one layer of lecithin are separated by the “tails” of a second layer facing the other way, as shown in Figure 5.14), either of which will hinder flow. The actual amount of lecithin that can be used before thickening occurs depends to a certain extent on the particle size distribution. A finely milled chocolate, with a large specific

\textbf{Figure 5.12} Confocal laser scanning microscope picture of lecithin (fluorescing) surrounding solid particles within the chocolate.
Figure 5.13 Influence of soya lecithin on the viscosity parameters of two dark chocolates: (1) 33.5% fat; (2) 39.5% fat. (Chevalley²).

Figure 5.14 Schematic diagram of a spherical micelle and a bilayer of lecithin on a sugar particle.
surface area, will have a relatively high yield value, as was explained earlier. This can be partially offset, however, by the fact that as there is a bigger area to cover, more lecithin can be used before this increase in yield value takes place.

At one time, for a product to be called “chocolate” on its label in a shop, the amount of lecithin used was restricted to 0.5% or 1.0% depending on the type of chocolate that was being made and where it was being manufactured or sold. The current EU regulations have changed this so that lecithins may be used according to the principle of *quantum satis* (no maximum level is specified). However, they must be used with the Good Manufacturing Practice directive and at levels no higher than those needed for the intended purpose. There is also a very small amount of lecithin which is present naturally in the cocoa and the milk components, especially in buttermilk.

Soya lecithin is a mixture of natural phosphoglycerides (phospholipids) with other substances such as soya oil (see Table 5.1). It is used widely throughout the food industry. Its composition, however, can vary, and some lecithin manufacturers have tried to optimise those components that are beneficial for chocolate flow, thus fractionated lecithins are commercially available. The phosphatidylcholine part of lecithin has been shown to be particularly effective in reducing the plastic viscosity of some dark chocolates (Figure 5.15), whereas other fractions have been shown to have a negative effect particularly on the yield value. Because the ratio of the different components varies within standard lecithins, its effectiveness in reducing chocolate viscosity can change from batch to batch. For this reason some suppliers provide a standardised product.

In order to try to overcome the increase in yield value at higher levels, Cadbury developed an alternative surface active agent from

<table>
<thead>
<tr>
<th>Table 5.1 Phosphoglyceride (phospholipid) composition of soybean lecithin.</th>
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<tbody>
<tr>
<td>Phosphatidylcholine (PC)</td>
</tr>
<tr>
<td>Phosphatidylethanolamine (PE)</td>
</tr>
<tr>
<td>Phosphatidylinositol (PI)</td>
</tr>
<tr>
<td>Phosphatidic acid (PA)</td>
</tr>
<tr>
<td>Other phosphoglycerides</td>
</tr>
</tbody>
</table>

The remaining approx. 44% is mainly triglycerides.
hardened rapeseed oil. This is an ammonium phosphatide and is known as YN.

Commercial lecithin is primarily manufactured from soya and, although lecithin is generally regarded as being beneficial to health, some countries have expressed concern that part of the soya may be derived from a genetically modified source. This has led to lecithin being sold as being from other sources or from selected growing areas such as Brazil. In addition, alternatives to lecithin are being marketed for confectionery use, e.g. citric acid esters. Their actual effectiveness is likely to depend on the chocolate type and the method of manufacture used to produce it.

### 5.5.2 Polyglycerol Polyricinoleate

Polyglycerol polricinoleate (PGPR), also known as Admul-WOL, is a very different surface active agent/emulsifier. It was originally developed for use in the baking industry and can be manufactured by the polycondensation of castor oil and of glycerol, the products of which are mixed and then esterified. Although having a relatively small effect on the plastic viscosity, it has a dramatic one on the yield value. At only 0.2% in the chocolate it can halve the yield value relative to lecithin, and at about 0.8% it has been shown to
reduce it to zero, turning the chocolate into a Newtonian liquid. How it does this has been the subject of a lot of studies, but is still not fully understood. Its effect is very significant and may be useful or undesirable. Figure 5.16 shows two chocolates pouring onto a biscuit, with the same plastic viscosity, but very different yield values due to the presence in one of some PGPR. Very often the required flow properties lie somewhere between these two extremes, and mixtures of PGPR and lecithin are commonly used by many chocolate manufacturers and indeed sold as such by commercial emulsifier suppliers.

5.5.3 Other Emulsifiers

Other emulsifiers such as sorbitan esters, spans and tweens etc. are often found in chocolates and chocolate flavoured compound coatings. They are normally less effective in reducing the yield value or the plastic viscosity than either lecithin or PGPR. They often do, however, alter the setting rate, the gloss on the product and especially its shelf life with respect to bloom formation. This will be described in more detail in Chapter 6.

5.6 DEGREE OF MIXING

The degree of shear/mixing is critical in order to obtain the thinnest possible chocolate for a given fat content. This to a large extent is
governed by the design of the conche and its mixing elements. There are other factors that are important, two of which are the order of addition of the ingredients into the conche and the drive motors to these mixing elements.

The conche is trying to coat the solid particles with fat and a smearing action does this very well, provided the particles are unable to move out of the way. If there is a lot of fat present they can move more easily and so the process becomes less efficient. This means that it is necessary to add only enough fat for the chocolate to turn into a paste. The remainder of the fat should be added towards the end in the liquid conching stage. It is said that fat added at this stage is twice as effective in reducing the final chocolate viscosity as the same amount of fat added at the beginning of conching.

It is even more important to add the majority of the lecithin at the later stages. One end of the lecithin molecule is very hygroscopic and will bind to water, making it more difficult to remove. This means that it should normally be added at the end of the dry conching stage, when a majority of the water has been removed and, preferably, like the fat, it should be added at the liquid conching stage. In certain circumstances a fraction of the lecithin is added at the beginning of conching to reduce the rate of moisture removal. If the conche is hot, or a lot of moisture is present, then the water will be released from the ingredients at a faster rate than it can escape from the conche. It then gets back into the chocolate and forms hard agglomerates by sticking sugar particles together, which makes the chocolate taste gritty. This can be alleviated by this early addition of some of the lecithin.

Some people believe, however, that if lecithin is present at the roll refining stage, then the pressure will force it into the cocoa particles making it ineffective. Other sources state that high temperatures will reduce the effectiveness of lecithin, but this also has never been fully substantiated.

How the mixing elements are controlled will also effect the viscosity of the final liquid chocolate. Traditionally the mixing arms had one or two speeds, and could be reversed once the chocolate became thin so that the wedge end cut into the liquid (as described in Chapter 4). If the power used by the conche is recorded against time, this gives a jagged curve similar to the one
illustrated in Figure 5.17a. For long periods, therefore, the conche is using very little power and therefore having very little mixing effect. Now it is possible to electronically control the process so that when the power drops, the speed is increased to bring it back up again. This means that a power curve like the one shown in Figure 5.17b is recorded. This type of control enables a thinner chocolate to be produced in a shorter time than when the traditional method was used.

Having produced the liquid chocolate, it is necessary to have the chocolate in a form in which it is still liquid, but will set quickly in the correct crystalline form to give the correct snap and gloss. This depends on the fat type (Chapter 6) and the way the chocolate is cooled and pre-crystallised (Chapter 7).

REFERENCES


CHAPTER 6
Crystallising the Fat in Chocolate

In order for a product to be sold as chocolate most of the fat inside it must be cocoa butter. This is a fat made up of several different triacylglycerols (triglycerides), each of which will solidify at a different temperature and at a different rate. To make it even more complicated, there are six different ways that the individual crystals can pack together. What makes it difficult for the chocolate maker is that only one of these six forms will give the product the good gloss and snap on breaking that makes it so attractive to the purchaser.

Milk chocolate must also contain milk fat. This will alter how it sets and also the texture of the final product. In some countries chocolate may contain a non-cocoa vegetable fat. When two or three fats are mixed together, the setting properties and texture are not a simple average of its components, as a phenomena known as fat eutectics takes place. This means that there is a limit to the number of types of vegetable fat that can be used in chocolate.

If the wrong type of fat is present or if the chocolate is old or has not been crystallised properly, then a white powdery surface forms, known as fat bloom. This is in fact made up of fat crystals and not mould. Special fats and emulsifiers have been developed to retard its formation and to make chocolate more able to withstand higher temperatures. Chocolates that have been left in the sun will bloom very rapidly.

Some confectionery products contain a chocolate flavoured compound coating. There are two main types of these, one which contains some cocoa butter/liquor and another which has only cocoa powder.
6.1 STRUCTURE OF COCOA BUTTER

All fats are mixtures of triglycerides, i.e. they have three fatty acids attached to a glycerol backbone. In cocoa butter there are three main acids which account for over 95% of those present. Almost 35% is oleic acid (C18:1), about 34% is stearic acid (C18:0) and approximately 26% is palmitic acid (C16:0). It is, in fact, because this fat is relatively simple in having so few main components that it melts rapidly over such a small temperature range, i.e. between room and mouth temperatures.

These acids are attached to the glycerol in a way that is illustrated diagrammatically in Figure 6.1. This shows palmitic acid (P) in position 1, oleic acid (O) in position 2 and stearic (St) in position 3. This is known as a POSt molecule. If the stearic and oleic acids were reversed this would become PStO, which is quite a different molecule even though the constituents are the same.

The stearic and palmitic acids are saturated acids, i.e. the hydrocarbon chain which makes up the fat does not contain any double bonds. In unsaturated fats this chain contains one or more double bonds, as is the case for oleic acid. The molecule in Figure 6.1 can therefore be described as symmetrical and mono-unsaturated and is often referred to as an SOS triglyceride, where S refers to any saturated acid. About 80% of cocoa butter is of this form, i.e. it has oleic as the middle acid.

Between about 1% and 2% of cocoa butter is all saturated (SSS, long chain trisaturated triglycerides, where the saturated fat is mainly palmitic or stearic) and melts at a much higher temperature.

Figure 6.1 Structure of triglycerides. Nominally this is the POSt molecule in the β form with all carbon zigzags similarly oriented and lying approximately in the plane of the molecule. Reprinted with permission of Loders Croklaan.
than the more common SOS. From 5% to almost 20% on the other hand contain two oleic acid molecules and are SOO, which is mainly liquid at room temperature. When these are combined, as they are in cocoa butter, the fat will therefore be partly liquid at room temperature. If milk fat is present this proportion will increase and the chocolate will be softer to bite into. As the temperature rises the fat will melt according to the proportions of the different types of fat present.

It is possible to measure the proportion of solid fat present at any temperature using techniques such as nuclear magnetic resonance (NMR, see Chapter 8) and the curve produced is known as the solid fat index. Figure 6.2 shows examples of this for cocoa butters from three different countries: Brazil, Ghana and Malaysia. In Chapter 2 it was stated that normally the nearer the equator the cocoa was grown the harder will be the fat. This is demonstrated clearly here, as at 20°C 81% of the Malaysian cocoa butter is solid compared with only 66% for the Brazilian one, with the Ghanaian one being in between. At 32.5°C the difference is proportionately

![Figure 6.2](image_url) Typical solid fat content of cocoa butters from Brazil, West Africa and Malaysia obtained by NMR measurements.
greater with only 7% of the Brazilian sample still being solid compared with 20% for the Malaysian one. It is possible to see the reason for this by looking at the triglyceride content of the different cocoa butters in terms of their saturated and unsaturated fats, given in Table 6.1. The relative hardness is particularly affected by the SOS/SOO ratio. This is 3.6 for Brazil but 16.5 for Malaysia.

When the cocoa butter is melted and then cooled, the three types of triglyceride once again behave differently. This is illustrated in Figure 6.3. The SSS types crystallise first. These make the chocolate thicker as there is less liquid fat present, but it is the SOS crystals

<table>
<thead>
<tr>
<th>Triglyceride</th>
<th>Brazil</th>
<th>Ghana</th>
<th>Malaysia</th>
</tr>
</thead>
<tbody>
<tr>
<td>SSS&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.0</td>
<td>1.4</td>
<td>2.3</td>
</tr>
<tr>
<td>SOS</td>
<td>63.7</td>
<td>76.8</td>
<td>84.0</td>
</tr>
<tr>
<td>SSO</td>
<td>0.5</td>
<td>0.4</td>
<td>0.5</td>
</tr>
<tr>
<td>SLS</td>
<td>8.9</td>
<td>6.9</td>
<td>6.8</td>
</tr>
<tr>
<td>SOO</td>
<td>17.9</td>
<td>8.4</td>
<td>5.1</td>
</tr>
<tr>
<td>OOO</td>
<td>8.0</td>
<td>6.1</td>
<td>1.3</td>
</tr>
</tbody>
</table>

<sup>a</sup> S = saturated fatty acids (mainly palmitic and stearic), O = oleic acid, L = linoleic acid.

Figure 6.3 Crystallisation temperatures and rates for the different triglyceride groups. (Talbot<sup>1</sup>)
that form later on that determine the chocolate’s texture and resistance to fat bloom.

6.2 DIFFERENT CRYSTALLINE FORMS

It is commonly known that carbon can exist in different forms, ranging from soft graphite through intermediate hardness materials, up to the very hard structure of diamond. Fats can also crystallise in a number of different ways, a property that is known as polymorphism. As the structure becomes denser and lower energy, it becomes more stable and harder to melt.

The reason for this is that the different fat molecules, as illustrated in Figure 6.1, can fit together in a number of different ways. Because of their shape, it is in a way like stacking chairs. There are two ways that this can be done, as illustrated in Figure 6.4, *i.e.* double-chain packing and triple-chain packing. These small stacks have then got to fit together with other stacks. The angle at which they fit together determines their stability. If chairs are stacked straight-up they tend to fall over. This is the same for fats in that a straight-up arrangement, known as alpha is formed at low temperatures and transforms rapidly into one of the other forms (see Figure 6.5).

The analogy with chairs then breaks down. The angle of tilt at which the crystals fit together then determines their stability, whereas chairs stacked at an angle would just fall over. Some fats have only one stable form whereas others have three, namely α, β' and β. Cocoa butter has six however. There are two nomenclatures that are used to describe these polymorphs. The chocolate industry tends to number them I to VI, as was described by Wille and Lutton in 1966. The oils and fats industry in general prefer the Greek letters defined by Larsson in the same year. Figure 6.6 shows the temperature ranges at which the different forms crystallise.

![Double- and triple-chain length packing configuration](image-url)
Figure 6.5 Crystal packing of tryglycerides. (1) Projection showing arrangement of alkyl chains for α, β, and β' polymorphs. (2) Projection looking onto the ends of chains. Reprinted with permission of Loders Croklaan.

Figure 6.6 Temperature ranges for the stable formation of the six different crystalline forms of cocoa butter.
Forms V and VI are the most stable and are triple-chain packing, whereas the other forms are all double.

More recent research by Van Malssen et al.\textsuperscript{2}, however, found only five polymorphic forms in cocoa butter. The main difference being that they found not two $\beta'$ forms but a range of these, indicating the presence of either five forms (if the range is treated as a single form) or many forms (treating the forms individually). This work also suggested that the melting points of each of the polymorphic forms had been wrongly assigned by Wille and Lutton and that the melting point of the lowest stability form (Form I) was significantly lower at $-5\,^\circ\text{C}$ to $5\,^\circ\text{C}$. This is possibly because this form transforms very quickly into Form II, making determination of its melting point very difficult. Because it is widely used within the confectionery industry and it explains most phenomena regarding crystallisation, the traditional Forms I to VI nomenclature is used in this chapter.

Form I is very unstable and melts at about $17\,^\circ\text{C}$ and so is present only on coatings for ice creams. It rapidly changes into Form II, which in turn transforms, although at a slower rate, to Forms III and IV.

If liquid chocolate at about $30\,^\circ\text{C}$ is used to make a typical confectionery product and is then cooled in an airflow at about $13\,^\circ\text{C}$ for about a quarter of an hour, Form IV will be the main crystal type present – unless some sort of pre-seeding has taken place. Form IV is relatively soft and so the chocolate will not have any snap when it is broken. In addition, it will transform over a period of days to Form V. The actual time depends on the storage conditions, with the transformation taking place faster at higher temperatures. The more stable forms are however more dense, so the chocolate will contract. Some of the cocoa butter is still liquid even at room temperature however, and, in addition to this, some energy is given out as the fat is transforming to a lower energy state. This combination of effects pushes some of the fat between the solid particles and onto the surface. Here it forms large crystals that give the white appearance of chocolate bloom, as shown in Figure 6.7.

For this reason it is necessary for the chocolate maker to ensure that the cocoa butter is in Form V when it is used to make confectionery products. This form is hard with good snap, gives a glossy appearance and has a relatively good resistance to bloom.
It will also contract well if the liquid chocolate is poured into a mould (see Chapter 7).

Form VI is in fact more stable, but under normal conditions, it is only formed by a solid to solid transformation and not directly from liquid cocoa butter. This means that chocolate with fat in Form V will, after a period of months or sometimes years, start to bloom. This is because the same effects as were described for the IV to V transition are happening again, but at a slower rate. Therefore the skill of the chocolate maker is to get the chocolate as rapidly as possible into Form V, but then take precautions to prevent the further transformation.

6.3 PRE-CRYSTALLISATION OR TEMPERING

If chocolate is cooled to 34°C and then stirred slowly, Form V crystals will eventually appear and, after a long time, probably days, there will be enough to seed all the remaining chocolate. This means that if it is cooled suddenly there are enough crystals present to form nuclei around which the remaining fat will set with the same type of crystal. This is obviously impractical for the chocolate industry, where several tonnes of chocolate an hour are often used.

Figure 6.7 Bloomed and unbloomed dark chocolate sweets.
Some other form of pre-crystallising, commonly known as tempering, is required. For small batches it is possible to add small amounts of previously set chocolate. Here a few per cent of grated solid chocolate is added to liquid chocolate which has been previously cooled to about 30°C. This is often described in cooking recipes for the home, when cocoa butter containing chocolates are being used (see Project 9 in Chapter 12). (Many cooking chocolates are in fact compounds which contain other fats. These fats solidify in only one crystalline form so seeding or tempering is not required). Recently a method has been developed to produce small cocoa butter crystals by spray chilling. Once they have transformed to Form VI, they are used to seed chocolate.

The speed that the fat in chocolate begins to crystallise depends not only on the temperature but also on the rate at which it is mixed and sheared. This is due to the fact that fat will solidify onto any seed crystal that is present. The seed therefore needs to be of the correct type and to be well distributed throughout the chocolate. Large crystals will have much less effect than the same amount of solid fat in small crystals that are uniformly mixed throughout the chocolate. High shear has the effect of breaking the solid fat crystals and uniformly distributing them. In addition, it provides heat and energy which increase the rate in which the more unstable crystals can change to Form V. Ziegleder\textsuperscript{3} showed that the effect was extremely dramatic and some of his results are given in Figure 6.8. This shows that by using extremely high shear rates cocoa butter can be pre-crystallised in 30 seconds rather than several hours or days.

There is a major problem with high shear rates however, in that they can generate too much heat, which will help the transformation to Form V. If there is too much heat, on the other hand, it will melt all the crystals completely.

The actual process used by chocolate manufacturers is a compromise that uses an intermediate shear rate. In order to increase the rate of crystallisation, the chocolate is cooled to temperatures where Forms II and III are created. It is thoroughly sheared to form a lot of smaller crystals, before being reheated, still under shear, to convert the unstable ones to the required form. The machines that do this are called tempering machines and further details about them are given in the next chapter. The actual temperatures used depend very strongly on the fats present in the
chocolate. These are mainly cocoa butter and milk fat, but in some chocolates non-cocoa vegetable fats are also present.

### 6.4 MIXING DIFFERENT FATS (FAT EUTECTICS)

When two or more fats are mixed together it is important that the final chocolate sets at a suitable rate and, more importantly, that it has the correct texture and melting properties in the mouth. One way to determine this is to measure the solid fat index, as was demonstrated for different cocoa butters in Figure 6.2. If instead of

![Figure 6.8 Relationship between crystallisation and the rate of shear (mixing) at different temperatures. (Ziegleder³)](image)
measuring the proportion of solid fat at different temperatures, the proportion is determined for different blends of two fats at the same temperature; then a plot is obtained as illustrated in Figure 6.9.

It might be expected that the solid fat index could be calculated from the solid fat measurement for the individual fats according to the proportion that they are present in the mixture, i.e. the straight line on the graph. As can be seen, this is not the case.

The reason for this is that although other fats, like milk fat, are triglycerides, their structure is very different from cocoa butter. Returning to the analogy of stacking chairs, it is as if chairs of another size are placed within the stack. This will make the overall structure far less stable. This means the product will melt more easily, i.e. it will contain far more liquid fat and this is what in fact happens for most fat mixtures, as shown in Figure 6.9.

If only a small amount of the other fat is present, the disruption it causes is less, so the actual hardness is near the expected one. Where the two fats are in similar proportions, the softening effect is largest. The degree of this difference and also the amount that can be added without causing a significant texture difference will depend on how differently the fats crystallise. As will be seen later,

Figure 6.9 Expected and the actual solid fat content at 20°C of mixtures of cocoa butter and a soft filling fat.
this is very important when adding vegetable fats, and very few can in fact be used.

Milk fat is present in all milk chocolates and even many plain ones. In the second case the reason is to reduce the possibility of fat bloom. If milk fat is added at about 5% of the weight of the chocolate, which contains a total of 30% fat, then it makes up about 17% of the fat phase. This makes it softer and also increases the time that it takes for the cocoa butter to transform from Form V to Form VI and produce a white coating on the surface.

In milk chocolate even higher levels of milk fat are sometimes used, but there is a limit, which is set by the softening (eutectic) effect. Chocolate eaters expect milk chocolate to be softer than dark chocolate, but they do not want it to be too soft. This once again can be determined from the solid fat index. Figure 6.10 shows data for milk fat/cocoa butter mixtures at different temperatures. The lines are drawn through points of equal hardness/liquid fat content. Most consumers eat chocolate at around 20 °C, and for it to be acceptable it must have about 70% solid fat. This means that about 15% milk fat is an upper limit. This measurement is, of course, for fat that is freely mixed with the cocoa butter. In looking

Figure 6.10 Melting profiles of fat blends of cocoa butter and milk fat. Reprinted with permission of Loders Croklaan.
at the different types of milk powder in Chapter 2, it was noted that some milk fat can be retained within the structure of the powder. This was said to be detrimental to the flow properties of the liquid chocolate, which it is. On the other hand, any bound fat does not contribute to the softening effects. Chocolates containing some bound fat have been manufactured with the milk fat accounting for over 27% of the fat phase.

The solid fat index not only tells us about the hardness of the chocolate, but also gives information about what happens when it melts in the mouth. Figure 6.11 illustrates curves for three different fat mixtures. The percentage of solids at room temperature gives an indication of the hardness of the chocolate. In this case, A is the hardest and C the softest. The temperature at which the sudden

Figure 6.11 Solid fat indices of three different fat mixtures.
reduction in solid fat content occurs relates to its heat resistance. Malaysian cocoa butter would normally be at a higher temperature than Brazilian butter (Figure 6.2). This makes it better for use in hot climates, but worse for ice cream. In this case, once again A melts at a higher temperature.

The steepness of the curve relates to how quickly it will melt. To change from a solid to a liquid requires a lot of energy. This is known as latent heat and for cocoa butter is about 157 J g\(^{-1}\). This can be compared with the specific heat of 2.0 J g\(^{-1}\), which is the amount of energy required to raise the temperature of the fat by 1 °C. When chocolate melts in the mouth, therefore, a lot of energy is required, which can only come from the mouth itself. (The temperature is raised by approximately 20 °C and it also melts the chocolate. The combined specific and latent heat is 20 \(\times\) 2 + 157 = 197 J.) If the slope is steep this happens very quickly and the mouth feels cool. Fats have been developed with exceptionally sharp melting curves, which exploits this property. The speed of melting also alters the viscosity of the chocolate before it is swallowed, which in turn changes the speed that the particles can reach the flavour receptors. So the flavour is also changed. In Figure 6.11, A and C will have similar cooling effects, both of which are greater than B.

Some fats, like B, have a significant amount of solid crystals still present even at 40 °C. This will not melt in the mouth and will leave a waxiness in the mouth. Some fats used to make chocolate flavoured coatings suffer from this problem. Other fats can be added which will reduce the amount of these high melting point fats by the eutectic effect. Care must be taken however as this can lead to chocolate bloom formation.

### 6.5 CHOCOLATE FAT BLOOM

There are four main ways that chocolate fat bloom is formed. Two have already been described, i.e. by the Form IV to Form V transformation following incorrect pre-crystallisation (tempering), or by the age and temperature related Form V to Form VI change, which can be slowed down by the addition of milk fat.

Another way is where the chocolate has been melted and recrystallises without retempering, e.g. when it has been placed in the sun. This can be overcome by adding fat crystals of the same shape as
cocoa butter, but with a very much higher melting point, to the chocolate. Unless these crystals are melted, which may require temperatures above 50 °C, they remain in it and will seed it when it sets again, preventing it from blooming. One of this type of high melting point seeding fat is made with behenic acid to replace the saturated acids on the glycerol chain. Behenic acid (C22:0) is, in fact, present in cocoa butter, but at much less than 1%. BOB (1,3-behenoyl-2-oleoylglycerol) is manufactured by the Fuji Oil Company in Japan, but cannot yet be legally used in many other countries of the world.

The fourth mechanism is due to the migration of soft fats into the chocolate. In a box of chocolates that contains a wide variety of centres, it is usually the nut centres that turn white first. Nuts like hazelnuts contain a fat that is almost all liquid at room temperature. Pralines are made similarly to chocolate but with hazelnuts. If the two are put together to form a sweet, the fat phase tries to reach an equilibrium (see Figure 6.12). The liquid fat from the centre moves into the chocolate and, because of the eutectic effect, makes it much softer and may even cause some of the cocoa butter to become liquid again, if the softening is bad enough. Some of the cocoa butter migrates in the reverse direction into the centre making that harder. Thus the difference in texture is lost and the sweet is less appealing. As the soft fat migrates to the surface, it

Figure 6.12 Migration of fats within a chocolate with a nut-based filling.
takes some of the cocoa butter with it. This crystallises forming fat bloom. It is also possible that this type of fat causes a rapid increase in the rate of the Form V to Form VI transition of the cocoa butter, which would also cause it to bloom.

There are several ways of reducing this type of bloom. Some are based on trying to stop the soft fat getting into the chocolate at all. This is done by having another harder fat that either forms a layer inside the chocolate shell or forms a sponge-like system within the centre. Alternatively an anti-bloom fat is added to the chocolate or to the centre, from where it migrates together with the soft nut oil. These fats have the property that they can slow down the Form V to Form VI transition, which would cause the fat bloom.

These are just some of the vegetable fats that can be used in confectionery. Others may be added to increase the heat resistance or to remove the need for tempering.

6.6 SOME TYPES OF NON-COCOA VEGETABLE FAT

Vegetable fats have been used in chocolate and chocolate-like coatings for many years. In the First World War, companies like Rowntree used vegetable fats in their chocolate because they were unable to buy cocoa butter. In the 1950s research showed that, unlike animal fats, some vegetable fats contained the same triglycerides as cocoa butter. This led to a 1956 Unilever patent, which showed a method of producing a fat that was almost identical to cocoa butter, but was made from fat obtained from other vegetable sources. These were made commercially and added to chocolate in different amounts. Current EU legislation restricts their use to 5% of specific fats, if the product is being sold as chocolate, and also requires very clear labelling. If higher levels, or other fats are used, it must be sold under another name, such as a chocolate flavoured coating. Legislation varies elsewhere in the world with a few countries even allowing all the cocoa butter to be replaced by other fats.

The original vegetable fat made by Unilever and the many others that are now on the market are known as cocoa butter equivalents (CBEs), as they are like cocoa butter and can be added in any proportion without causing a significant softening or hardening.
effect. Other fats can be used only if almost all the cocoa butter is replaced, and these are known as cocoa butter replacers (CBRs).

6.6.1 Cocoa Butter Equivalents

To be able to be added to cocoa butter without having a eutectic effect, the vegetable fat must crystallise in the same way as cocoa butter (i.e. using the chairs analogy, have the same size and shape chair). Cocoa butter contains palmitic (P), stearic (S) and oleic (O) acids on a glycerol backbone, with the majority of the molecules being POP, POS and StOS. The fat manufacturer has therefore to obtain these different fractions from different sources and then blend them.

The POP is the easiest to find as it is a major component of palm oil, which is obtained from the palm (*Eleaeis guineensis*) widely grown in Malaysia. A lot of other fats are present, but these can be removed by fractionation. The easily melting (olein) and the hardest melting (stearin) parts are removed, leaving the mid-fraction, which is mainly POP and a small amount of POS. Two types of fractionation are used: dry fractionation and solvent fractionation. In dry fractionation the fat is heated to a predefined temperature and then the liquid part separated from the solid by pressing or filtering. In solvent fractionation, the fat is dissolved normally in acetone or hexane. The higher melting triglycerides are then allowed to crystallise and are filtered out. This procedure gives much more clearly defined fractions than the dry process.

The StOS and increased amounts of POS are much harder to obtain. One nut that gives this type of fat is illipe (*Shorea stenoaptra*). This is grown in Borneo, but is only intermittently available. Shea (*Butyrospermum parkii*), which grows in West Africa, and sal (*Shorea robusta*), from India, both contain a high proportion of StOS, but the crops are not always available and can be of poor quality. However, by blending the palm oil mid-fraction with illipe fat and the harder (stearin) fraction from shea fat, it is possible to manufacture a fat that is totally compatible with cocoa butter.

In summer in southern Europe or in tropical climates, the chocolate will melt easily under ambient conditions. By altering the proportions of StOS it is possible to make the chocolate so that it will not melt until the temperature is several degrees higher than
normal cocoa butter, but does not leave a waxiness in the mouth. If the solid fat index were measured as in Figure 6.11, the nearly vertical melting curve would move to the right and there would be no significant amount of fat that was solid above about 36 °C. This type of fat is known as a cocoa butter improver (CBI), because it improves the chocolate. The harder fat components are difficult to obtain, so CBIs are more expensive than cocoa butter equivalents.

### 6.6.2 Enzyme Interesteralification

Because good quality harder fats are difficult to obtain, fat manufacturers have developed a process that enables them to use other raw materials such as sunflower oil. This is used to obtain a fat that can be blended with palm oil to give the required cocoa butter equivalent.

Enzymes occur naturally, both inside and outside the human body, and have the ability of being able to speed up the separation of the fatty acids from the glycerol backbone leaving partial glycerides (monoglycerides and diglycerides) as well as glycerol. One of these, called lipase, occurs in flour and can give important problems to the confectionery manufacturer because, with certain fats, it produces free fatty acids that have a very unpleasant cheesy flavour. Some of these enzymes are position specific and will accelerate only that reaction which cuts off acids from specific positions on the glycerol. They are also able to exchange the fatty acids in certain positions as well as remove them. The fat manufacturers use this property to change the melting properties of their fats.

In the 1970s workers at Unilever Research Laboratory at Colworth House in England showed that a certain type of enzyme, called mucor miehei, attacked only the 1 and 3 positions of a triglyceride. The process they developed can use any fat that mainly has oleic acid in the 2 position (see Figure 6.13). This is then mixed

![Diagrammatic representation of enzyme interesteralification.](image-url)
with stearic acid and the appropriate enzyme. This free stearic acid is then esterified onto the 1 and 3 positions, where the original fats are removed by the enzyme. The central oleic acid remains untouched, so the fat becomes very rich in StOSt. The free fatty acids released by the enzymes are then treated by de-acidification and then fractionation and refining remove the impurities. This fat is identical to that obtained directly from other tropical nuts in terms of its processing to make confectionery and its behaviour in the mouth when eaten. It can also be added to chocolate at any level that is required (although it cannot be called "chocolate" in the EU), as it is fully compatible with cocoa butter. Some other fats are very different, however.

6.6.3 Lauric Fat Cocoa Butter Replacers

There are other fats that melt in the same temperature range as cocoa butter, and so have a similar texture and mouthfeel, but which crystallise in a very different way. Palm kernel oil and coconut oil are both widely available and contain about 50% lauric acid (C12:0, dodecanoic acid), a lot of which exists as trilaurin. Unlike cocoa butter it sets in one crystal form and so no pre-crystallisation is required. This form is also different in that it is not a β (like Form V in cocoa butter) type crystal but a β′ one (equivalent to Form IV).

Once again we are in the situation of trying to stack different chairs. This is demonstrated clearly by measuring the solid fat index of mixtures of a lauric fat with cocoa butter, as shown in Figure 6.14. Apart from where only about 5% of the other fat is present, this mixed crystal system exists. It is very soft, will take a long time to solidify and is likely to bloom at a very rapid rate.

This means that these lauric cocoa butter replacers can be used only when very little cocoa butter is present. Because cocoa liquor contains about 55% cocoa butter, the products made with lauric cocoa butter replacers are normally made with cocoa powder. This in turn tends to give them a different flavour. They are however often made into coatings because, for the small confectioner or home user, it is a big advantage not to have to temper it. Cooling this type of fat must be very rapid and the products are often initially very glossy.
With lauric fat products, however, it is very important to keep them in a dry environment and, if possible, ensure all other ingredients being used are lipase free. This is because in a moist environment the lipase enzymes will accelerate the removal of some of the free fatty acids from the glycerol backbone. Where lipase is present, the acids produced have a very unpleasant soapy flavour, even at a very low level in the product. Other fats do exist, however, which have slightly more compatibility with cocoa butter.

6.6.4 Non-Lauric Fat Cocoa Butter Replacers

Palm oil and soybean oil contain many of the same fatty acids as cocoa butter and these can be fractionated out leaving mainly the stearic, palmitic and oleic acids. These, however, are far more random than in cocoa butter, so, for instance, the oleic acid is very often in the 1 or 3 position. In addition, these oils often contain a significant amount of elaidic acid (C18.1, octadec-trans-9-enoic acid). This is an unsaturated fatty acid in the trans form. This means that the hydrogen atoms that are associated with the unsaturated carbon atoms lie on opposite sides of the double bond as illustrated in Figure 6.15 and results in its structure being very different from the unsaturated fat in cocoa butter – oleic acid.
(C18.1, octadec-\textit{cis}-9-enoic acid). The \textit{cis} form has the hydrogen on the same side of the bond, as also illustrated in Figure 6.15.

This more random structure coupled with this different unsaturated acid means that this type of fat has limited compatibility with cocoa butter. Measurements of the solid fat index for different mixtures are given in Figure 6.16. The graphs show once again the soft texture, conducive to bloom conditions occur when both fats are present in similar amounts. However the effect is less severe than for lauric fats. In addition, about 7\% of lauric fat can be added to cocoa butter, but, more importantly for the chocolate manufacturer, up to 25\% of cocoa butter can be present with a non-lauric cocoa butter replacer. This means that chocolate

\begin{figure}
\centering
\includegraphics[width=\textwidth]{ cis_trans_forms.png}
\caption{Illustration of \textit{cis} and \textit{trans} forms.}
\end{figure}

\begin{figure}
\centering
\includegraphics[width=\textwidth]{ solid_fat_content.png}
\caption{Solid fat content of a mixture of cocoa butter and a non-lauric CBR. Reprinted with permission of Loders Croklaan.}
\end{figure}
flavoured coatings can be made with cocoa liquor, which in turn makes the flavour more like normal chocolate.

Non-lauric fats, like lauric ones, tend to set in the $\beta'$ form and do not need tempering, although it may be better to do so if a significant amount of cocoa butter is present. They also need much slower cooling than the lauric fat cocoa butter replacers.

### 6.6.5 Low Calorie Fats

In the USA in particular, there has been a demand for low calorie products. Normal fat contains 9 kcal g$^{-1}$ compared with 5 kcal g$^{-1}$ for the sugar and protein components. If the proportion of fat is reduced, the calorific value will fall. For processing and texture reasons, however, it is not possible to reduce the level too much below 25%. This is insufficient to make a low calorie claim on the product, so two manufacturers have produced fats that melt like cocoa butter but have a lower calorific value. Like lauric fats they are incompatible with cocoa butter and so the products have to be made with cocoa powder.

Proctor and Gamble produce a fat called Caprenin from caprylic (C8), capric (C10) and behenic (C22) acids. Caprylic and capric acids are metabolised in the body in a different way from the cocoa butter fatty acids, whilst, because of its chain length, behenic acid is poorly absorbed, if at all. This means that this fat is closer to a carbohydrate and has a declared calorific value of 5 kcal g$^{-1}$.

Salatrim (Benefat$^\text{TM}$) is produced by Nabisco and contains a mixture of long and very short chain fatty acid triglycerides. This too is said to have a calorific value of 5 kcal g$^{-1}$. Both these fats can only be used in a very limited number of countries.

### REFERENCES


CHAPTER 7

Manufacturing Chocolate Products

Having made the liquid chocolate it is then necessary to turn it into a solid bar, which may or may not have a centre of some other material, such as wafer, biscuit or fondant etc. First of all, however, it is essential to ensure that the fat sets in the correct crystal form using a procedure called tempering. A simple chocolate bar can then be made by pouring the tempered chocolate into a mould. Other moulded products have a chocolate shell surrounding a solid or semi-solid centre or, in the case of Easter eggs, this centre is left hollow. These products are produced by a process known as shell moulding.

In Roald Dahl’s famous story Charlie and the Chocolate Factory, there are chocolate waterfalls. These actually exist only in miniature, in a machine called an enrober. This produces a curtain of chocolate about 3 cm high which falls over the sweet centres as they pass through it on a wire belt. Many well-known products such as Mars Bar®, Lion Bar® and Crunchie® are made in this way.

A third way of coating centres is called panning. This is used to cover hard centres such as nuts and raisins with a chocolate coating.

Whichever form of processing is used, the chocolate must be allowed to set, so that it can be handled and packed. This must be done correctly, otherwise the chocolate will rapidly become white, with one of two forms of bloom.

7.1 TEMPERING

7.1.1 Liquid Chocolate Storage

This is the process which pre-crystallises a small amount of the fat in the chocolate, so that the crystals form nuclei, which help the fat set
rapidly in the correct form. The actual amount of fat that it is necessary to crystallise is uncertain, but is probably between 1% and 3%.

The liquid chocolate leaves the conche at a temperature normally above 40°C. If it is to be processed in the same factory, it is then kept in a storage tank until required. These tanks can hold 20 tonnes or even more and must be able to be stirred, heated and kept in a lower humidity environment.

If the chocolate is not stirred for a long period some of the fat separates on top, leaving a thicker chocolate in the bottom of the tank. The temperature is maintained at about 45°C. Keeping it for longer periods at higher temperatures would cause it to change its flavour, whilst the proteins in milk chocolate would tend to aggregate, making it become thicker. A much lower temperature would start the crystallisation process, and there would be the risk of the tank contents becoming solid. When describing the transport of cocoa beans it was noted that if the relative humidity was above the equilibrium relative humidity (ERH) of the beans, then they would take up moisture. This is equally true for the liquid chocolate. Here the ERH is between about 35% and 40%, so if the relative humidity around the storage tank is higher than this, the chocolate will take up water. Only a very small amount of water is required to start sticking the sugar particles together and increase the viscosity, making it difficult to process.

7.1.2 Tempering Machines

The tempering machine must first of all cool the chocolate down so that crystals can start to form. Chocolate is a very poor conductor of heat, so for it to cool quickly it must be mixed well so that all of it comes into contact with the cold metal surface of the tempering machine. These machines are a type of heat exchanger, which heats and cools the chocolate as it passes through it. A typical chocolate temperer is illustrated in Figure 7.1.

The centre column is a rotating shaft to which is attached a series of discs, or scrapers. The walls have bars or discs fitted to them, to ensure that the chocolate has to flow alternatively to the wall and to the centre and cannot just pass straight through the machine. The faster the shaft turns, the higher the shear rate and the faster the crystallisation. In many temperers this is between 3000 and
Figure 7.1 Schematic diagram of Sollich Solltemper MST-V tempering machine. (Nelson1)
8000 s$^{-1}$. The limit is set by the size of the motor and the heat developed by the mixing, which would start to melt the crystals.

The outside surface must be carefully temperature controlled and this is used to determine the degree of temper developed. The temperature is normally controlled within set groups of plates, called zones. Many machines have three or four zones, although many more are possible.

The first zone cools the chocolate to a temperature where crystals can start to form. The second takes it much lower so that Form IV and Form V crystals are also present, and during which the chocolate is highly sheared. The final stage raises the temperature to about 30 °C, which melts many of the unstable crystals.

The viscosity of chocolate, like most other substances, falls as the temperature rises. The thinner the chocolate, the easier it flows into a mould or around a sweet. It is therefore better to process the chocolate at as high a temperature as possible, without melting out the seed crystals. When they are newly formed, the crystals are small and are easily melted. If the chocolate is stirred and slowly heated, these crystals become more stable and have a higher melting point. It is for this reason that some tempering machines have an additional stage, where the chocolate is sheared more slowly and the crystals are allowed to “mature”.

### 7.1.3 Hand Tempering

For small-scale chocolate production, a tempering machine may be too large and hand tempering is used. This is performed on a marble table, which is heated to different temperatures in specific areas. The chocolate is poured onto one of the cooler areas, where it is mixed thoroughly using a scraper (Figure 7.2a and b). This thorough mixing and cooling initiates crystallisation and the chocolate is then moved to a warmer area of the table to melt the more unstable forms. This is very much an art and people who are skilled at it can then determine whether it is tempered by dabbing a small amount of chocolate on their lip. If it is tempered they are able to detect a defined cooling sensation. The less skilled operators use the method described below.

### 7.1.4 Temper Measurement

When the chocolate leaves the tempering machine it is important for the chocolate manufacturer to be sure that there are enough
crystals of the correct type to make the chocolate set properly. In the laboratory it is possible to use X-ray to determine exactly the types of crystal present. This is not only very expensive and slow, but the sugar is often removed first as its X-ray peaks would

Figure 7.2 Hand tempering; (a) pouring the chocolate; (b) scraping the chocolate.
dominate the spectrum. This may mean that the sample has changed by the time the measurement is taken, which makes it totally impractical for an industrial process.

The differential scanning calorimeter (DSC, see Chapter 8) is able to be used to provide a lot of information about the crystals present in the chocolate, but requires very careful sample preparation. This type of machine can be used for investigative work, but is too expensive and complicated for routine quality control.

In Chapter 6 in the section about solid fat indices, it was noted that fats sometimes give a cooling effect in the mouth because the energy required to melt them – the latent heat – is very much larger than the energy required to change its temperature by 1 °C – the specific heat. When solidifying chocolate this is also true, except that the latent heat is given out when the fat changes from a liquid to a solid, not taken in, as is the case in the mouth. The manufacturer can use this effect to determine whether the chocolate is correct, by measuring a cooling curve.

This can be done very simply and cheaply by the type of device shown in Figure 7.3. Typical cooling curves produced by this instrument are shown in Figure 7.4. The instrument is made up

Figure 7.3  A temper meter.
of a metal tube with a cup in the top to hold the chocolate sample. A thermometer probe fits through a cap, which fits tightly onto the cup. The temperature recorded by this probe can be recorded on a chart recorder as shown, or on a computer database. The tube fits through a holder in the thermos flask, which keeps the tube in position, with the cup above the water level in the flask.

Figure 7.4 Curves from temper meter for: (a) correctly tempered chocolate; (b) under-tempered chocolate; (c) over-tempered chocolate.
A mixture of ice and water is placed in the flask and then the tube is put in position. Some of the tempered chocolate is poured into the cup and the thermometer and cap pushed on top. A plot is then made of the temperature against time.

Initially the temperature drops uniformly but slowly. If the chocolate is tempered correctly, there is enough seed distributed throughout the chocolate to make it set very quickly. As it does so the latent heat is given out, and this offsets the cooling of the ice so the temperature remains constant for a significant time, as shown in Figure 7.4a.

If not enough seed crystals are present, it takes longer for the chocolate to set. There is however more fat to set and so more latent heat to come out. This means that there is a longer initial drop in temperature, but then so much latent heat is released that the temperature of the chocolate rises again. It eventually falls again once the latent heat has come out. This is shown in Figure 7.4b.

If almost no seed crystals are present, or indeed far too many (so that a lot of the latent heat has already been removed), a curve like Figure 7.4c is obtained, which means that the chocolate is not going to set correctly. Here mainly specific heat is being removed. The latent heat comes out slowly and, in the case of the uncrystralised chocolate, will be less because the more unstable crystals will be formed with their lower energy state.

The temperature at which the inflection on the curve occurs, point I on the curves in Figures 7.4a and b, is also very important. The higher the temperature, the more mature the crystallisation and the higher the temperature at which the chocolate can be used for moulding or enrobing.

There are several computer-based instruments on the market that calculate the rate of cooling and use this to determine factors such as one called a temper number, whilst others work in-line and can be used to automate the tempering process. All operate using the principle described above.

7.2 MOULDING

7.2.1 Solid Tablets

Moulding is the simplest method of forming chocolate and is what is used to make the simple tablets. For many years metal moulds
were used, but these were heavy, noisy and rather expensive. The latter is very important when a moulding line can contain as many as 1500 moulds, which must all be changed when the marketing department alter the weight or shape of a product. Currently most moulds are made of plastic, which is lighter and makes less noise. It also has the advantage that it can be twisted, which helps the solid bars to be removed when sticking occurs.

If the tempered chocolate comes in contact with a warm surface, the crystals within it will melt, so there won’t be enough seed for it to set properly. On the other hand, touching a cold surface may cause some of the fat to set in the wrong form. This will act as the wrong type of seed during subsequent cooling. It is therefore important that the empty moulds are pre-heated to within a few degrees of the temperature of the tempered chocolate before the start of the moulding process.

The chocolate is fed into the mould through a depositing head. This contains the same number of nozzles as the number of indents in the moulds below. A tablet line of width 1 m may have 10–20 nozzles. The moulds are transported below the depositor (a typical design is shown in Figure 7.5). The chains, or other transporting

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![Figure 7.5](image-url)  
**Figure 7.5** Diagram of piston-type depositor to put a measured amount of chocolate into a mould.
mechanism sometimes lifts up the moulds until they are just below the nozzles. A rod of liquid chocolate then falls along the indents as the moulds move forward and the chocolate continues to come through the nozzles. These depositors are designed so that an exact weight goes into each sector of the mould. As soon as this weight has been deposited, the mould quickly drops to its former level. This breaks any strands of chocolate which may still be attached to the nozzle.

The chocolate then needs to be spread evenly throughout the mould and any air bubbles must be removed to avoid blemishes of the type shown in Figure 7.6. This is done by vigorously vibrating the mould. In Chapter 6 chocolate was stated to be a non-Newtonian liquid in that it had a yield value. This means that energy must be applied to start it moving. This yield value will also tend to stop bubbles rising through the chocolate. Provided the shaker has the correct frequency and amplitude, however, it can turn a non-Newtonian liquid into a Newtonian one and therefore make the bubbles come out more easily.

When the liquid chocolate is at rest, the particles are practically in contact with one another and so it is difficult for them to move (Figure 7.7a). The vibration provides the energy to separate them, thereby lowering their resistance to movement and the yield value. When a bubble rises it does so very slowly, so we need to know what the viscosity appears to be at a low shear/flow rate. Figure 7.7b shows a plot of the apparent viscosity at a low flow rate against the frequency of the vibration for three different
amplitudes. This shows that we must vibrate this particular chocolate at more than 10 cycles s\(^{-1}\) and with an amplitude of at least 0.2 mm. If we use vibration, the chocolate has one seventh of its viscosity when it is vibrated at 30 cycles s\(^{-1}\) than when it is unvibrated. Low vibration rates do very little, so the manufacturer is wasting money in installing the vibrator and also on the energy involved. Very much higher rates do not improve efficiency. This thinning of the chocolate by vibration takes place only on the vibrator and stops as soon as the mould leaves it.

### 7.2.2 Chocolate Shells

Many moulded confectionery products contain a centre which contrasts with the chocolate in terms of taste or texture, e.g.
caramel, fondant or praline. Others like Easter eggs are just hollow shells. Both can be made on shell moulding lines. A schematic diagram of the procedures involved is given in Figure 7.8.

The first part of the process is the same as for making solid tablets. However, instead of letting the chocolate fully set, the moulds are passed through a short cooling tunnel so that only the outside sets. The mould is then turned upside down and vibrated. The chocolate initially sets near the mould wall, but should be still liquid in the centre. This liquid part runs out when the mould is inverted, leaving the outer shell.

There are several critical features to this procedure. Firstly the chocolate viscosity must be correct, in particular the yield value. If it is incorrect, for instance too high, none of the chocolate may come out. If it is too low, too much may come out and the shell will be too thin. Incorrect viscosities also lead to a non-uniform thickness of the shell. Thin sections may allow the centre material to leak from the final product (Figure 4.3), or to be very easily broken in the case of Easter eggs. Secondly the cooling period must be correct as it has to be long enough to let the shell form, but not so long that most sets, or that contraction occurs and the shell falls out too. As with the removal of bubbles from solid tablets, the
shaker frequency and amplitude must be correct in order to obtain good weight control of the shells.

Once the centre chocolate has run out, the moulds are turned back over so that the chocolate shell can be filled. In the case of hollow products, just the mould is further cooled so that the chocolate fully sets and contracts. The mould can then be turned over again and the shell will fall onto a conveyor belt ready for packing. Getting all the half shells for Easter eggs the same weight is not easy, so some packing lines are constructed such that the shells are weighed and divided into three types, heavy, average and light. They are then packed as two average halves or as a heavy one with a light one.

There are other ways of making hollow eggs and figures. These include book moulds and spinning. In the first technique, two moulds are hinged together and the shell moulding procedure carried out as normal. The rims are then melted and the hinge brought over like a book to bring the two parts together.

For spinning, the two halves of the mould are clamped together and a measured amount of tempered chocolate fed into it. The mould is put on a rotary arm, which turns and rotates so that all the inside is uniformly covered with the setting chocolate. Once it is hard enough, the mould is opened and the hollow figure removed.

When filling the shell, it is important that the centre material does not melt the chocolate. This is relatively easy for fondants or fat-based fillings such as pralines, which can be relatively fluid at about 30°C. With caramel this is normally much more difficult. Here the viscosity changes very rapidly with temperature (see for example Figure 7.9). It is necessary for the caramel to fill the shell and form a flat top, as any protruding strings or "tails" may penetrate through the chocolate base (see Figure 7.10). The chocolate normally acts as a moisture barrier to stop the centre drying out (or picking up moisture in the case of centres like wafers), but these tails provide a path for moisture transfer, which will shorten the shelf life of the product. In addition, any stickiness on the outside may cause problems during wrapping. This means that the caramel must be added as hot as possible, to make it flow as easily as possible and yet not so hot that it will melt the chocolate shell.

Once the centre is in place, then a chocolate base must be put on the shell. This is often done by having a scraper blade pressing
against the mould, as shown in Figure 7.11. Tempered chocolate is poured onto the moulds in front of the blade and forms a rotating roll in front of it. This fills in any remaining indents or holes, whilst the blade removes any excess chocolate.

Figure 7.9  Change in caramel viscosity with temperature.

Figure 7.10  A caramel “tail” within a chocolate-coated sweet.
The moulds are then cooled in a cooling tunnel, to set fully the shell and base, before they are turned over again and the sweets dropped onto a belt, which carries them into the packaging room. Because the outside fat has set in contact with the mould surface, this makes it smooth and very glossy. In any chocolate selection box, the moulded sweets are normally much shinier than those made using the enrobing process.

7.3 ENROBERS

In enrobing, the centre material, be it nougat, biscuit, fondant or caramel is first of all made separately and then placed on a belt, which will take it through the enrober. The aim is to get all the surfaces, including the base and ends, coated uniformly with chocolate. Typical examples that are made by this process are Mars Bar™, After Eight™ and Cadbury’s Crunchie™.

As with moulding, it is necessary to use tempered chocolate. The tempering unit may be located in the enrober line, or placed next to it with the chocolate being transferred in jacket-heated pipes.

A typical enrobing system is illustrated in Figure 7.12. The centres are placed on a continuous moving wire chain belt (A),
as shown in Figure 7.13, which transports them underneath the chocolate “waterfall” (B). Just beneath the belt at that position is the bottoming or surge trough (C). This retains the chocolate that falls through the chain belt, and recirculates it by means of the roller which is at the front end of the trough it. The chocolate from the
roller moves along with the centres and pushes them up slightly, forming a layer of chocolate on the undersides. Sometimes there are two “waterfalls” within a single enrober, at other times two enrober systems are used one after the other. This is particularly useful when the product has an uneven surface, e.g. crispies sticking out. The first coating will have a low yield value so that it flows into all the crevices and provides a good moisture barrier for the product. The second may have a high yield value so that the chocolate “stands up”, giving the product a more rugged appearance.

Having poured chocolate over the centres, it is then necessary to make sure that each item has the correct weight and appearance. This is done by a series of devices, whose effect can be varied according to the product being made.

The first device sends in a jet of warm air (D) which blows the excess chocolate from the top and sides of the sweet so that it passes through the belt and back into the base trough of the enrober (E). This tends to leave a ruffled wavy surface, especially if the chocolate has a high yield value. This is then smoothed, and excess chocolate is removed by the shakers (F). As was described earlier, these should be operated at the correct frequency and amplitude to overcome the yield value.

The next part deals with the bases. The chocolate was put on in the bottoming trough (C), but now may be too thick, or indeed be uneven and have uncovered areas. This is dealt with by the grid-licking rolls (G). Depending on the design of machine, these normally vary in number from one to three and may operate at different speeds and distances from the wire chain belt. This enables chocolate to be put on or removed from the base. Any remaining chocolate on the rolls is scraped off by a knife and once again falls back into the base trough.

The next section of the belt can be used to decorate the top of the sweets. For many years this was done by hand using marking forks of the type shown in Figure 7.14. Now most products are marked by a rotating roller, which just touches the top, or by programmable moving nozzles, which are able to put a wide variety of markings on the top in the same or even a different coloured chocolate.

Finally the product passes onto another belt, made of plastic, which passes through the cooling tunnel. This belt is often embossed with the name of the manufacturer or the product brand. As the
chocolate is still semi-liquid, the base then takes up this lettering. This is another way, other than the gloss, of determining whether a product has been moulded or enrobed, as an enrobed sweet will have any detailed lettering on the base, with less distinct top markings, whereas moulded sweets are able to have much greater detail on the top and sides, but the base is normally plain.

Frequently this plastic belt moves at a greater speed than the wire one. This separates the rows of product, making them easier to select for packing. It does, however, pull the sweet from the wire belt, often leaving long thin tails of the liquid chocolate behind it (see Figure 7.15). When these tails remain attached, the product is difficult to pack and often has to be sold as a misshape. In order to remove them, a small roller is placed between the two belts (H). This rotates at high speed and pulls the tails from the product. In order to work properly it must be correctly located. If it is too low it will have no effect, whereas too high a setting will mark the base of the sweets.

7.3.1 Maintaining Tempered Chocolate

Tempered chocolate is in a very unstable state. The temperature must be low enough not to melt out the seed crystals, which means
that other crystals will be forming. Eventually the chocolate will become thicker, and after a very long time it will solidify.

In the enrober the tempered chocolate is fed into the base trough (E). From here it is pumped up a pipe (I) into the flow pan (J). This pan is across the full width of the wire belt and has one or two slits along the bottom, which form the “waterfall”. As was described earlier, much of the chocolate from the flow pan passes through the wire belt due to blowing or shaking, or simply because it was poured over an area of the belt where no sweet was present. Eventually in the base trough chocolate would become thick and over-tempered.

Newly tempered chocolate is added to the base trough to replace that which is removed by the product, but normally this is not enough to stop this thickening occurring. In order to maintain stable conditions, therefore, some of the chocolate is continuously removed from the enrober base tank. It is then heated to between 40 °C and 50 °C to remove all the fat crystals, and then returned to the input of the tempering machine.

7.4 SOLIDIFYING THE CHOCOLATE

When the chocolates leave the moulding plant, most of the fat is still in the liquid state. In order for it to be firm enough to handle
and package, most of this must solidify in the correct crystalline form. This requires the removal of a large amount of latent heat and a relatively smaller amount of specific heat. The chocolates are probably already at a temperature several degrees lower than 30°C and, when they are ready for packing they will be at around room temperature, a drop of less than 10°C. The specific heat of chocolate is about 1.6 J g\(^{-1}\) C\(^{-1}\), so about 16 J must be removed from each gram of chocolate. The latent heat, on the other hand is 45 J g\(^{-1}\), so 45 J is removed for each gram, giving a total of just over 60 J to solidify and cool that amount of chocolate.

There are three ways that a body can lose heat, i.e. conduction, radiation and convection. In conduction the heat flows through the material directly in contact with that being cooled. In this case, the chocolate is in contact only with the plastic mould or belt. Both these are very poor conductors of heat, so very little heat energy can escape by this method, although a cold surface below the belt will help crystallise thin, flat products. Radiation heat transfer takes place at a rate determined by the fourth power of the temperature difference between the object being cooled and its surroundings (\(\Delta T^4\)). In this case the sweet is at about 25°C. If we assume that the surroundings are at 0°C, which, as will be shown later, is lower than may be desirable, then the heat transferred to this low temperature absorber will be 126 W m\(^{-2}\).

The third method is to blow cold air over the product. Some of the heat is given up to this air, which then moves away and is re-cooled. If the air is cooled to 0°C and blown over the product at 240 m minute\(^{-1}\), then heat can be transferred at 630 W m\(^{-2}\), i.e. more than five times faster than by radiation (Nelson\(^3\)).

Low temperatures, however, may give rise to two problems. Firstly they may cause the fat to set in the wrong crystalline form. This will cause the product to bloom very quickly and moulded chocolates not to contract properly and so become difficult to demould. Secondly moisture in the air may condense on cold surfaces and then drip onto the chocolate. This will dissolved some of the sugar from within it. When the chocolate is re-warmed ready for packing, the water evaporates again, leaving a white powdery surface. This looks very like fat bloom, but is in fact sugar and is therefore called sugar bloom. If possible, the temperature of the surrounding tunnel should be maintained above the dew point, i.e. that temperature at which moisture just starts to condense.
7.4.1 Coolers

Many coolers are just long tunnels, with blowers and coolers at intervals so that the tunnel can be divided into different temperature zones. A diagram of the cross section at such a position is given in Figure 7.16.

The initial cooling, particularly for enrobed products is fairly gentle. This is then followed by the coldest part, which corresponds to the position where most of the latent heat is given out. This is normally about 13°C, but lower temperatures are possible, provided the air is moving rapidly to prevent any condensation. The temperature is then raised slightly before the product enters the packing room. This is because if the product surface temperature is lower than the dew-point temperature in this room, moisture will condense on the chocolate causing blemishes, or maybe sugar bloom.

Where space is limited, and especially for moulded products, multi-tier coolers are used. Here the product goes backwards and forwards through different temperature zones. A three tier cooler, therefore, has a cooling time six times that of an equivalent length tunnel cooler.

The main problem with the multi-tier system is that the product must be kept horizontal as it is lifted from one level to the next one. This is sometimes achieved by a continuous chain driven system,
which holds the moulds or plastic trays on which enrobed sweets are placed.

The time it takes for a product to set depends not only on the quantity of crystals already in the fat, but also on the chocolate type and amount. A large block of chocolate is naturally going to take much longer than some small morsels being made for cookies. Normally, however, it takes between about 10 and 20 minutes to obtain a good quality product.

7.5 PANNING

Panning is used to make the small rounded chocolate items often sold in bags or tubes. These can be divided into two types: the chocolate-coated varieties, normally containing a nut or dried fruit and the chocolate-centred, sugar-coated products such as Smarties™ and M&M’s™. Both can be made using open rotating bowls or pans (Figure 7.17), often constructed of copper and which have temperature- and humidity-controlled air blowing into them through a pipe. These have often been replaced on the industrial scale by large rotating drums (Figure 7.18), which resemble very large washing machines and can hold more than 2 tonnes of

![Figure 7.17](image_url) An open pan for chocolate or sugar coating.
product at a time. These too have temperature- and humidity-controlled air and the facility to spray liquid ingredients onto the tumbling centres within the drum.

### 7.5.1 Chocolate Coating

For chocolate-coated products it is important to spend some time selecting good centres to coat with chocolate. It is better if all of them are of a similar size. As all the centres tumble over each other in the rotating pan or drum (Figure 7.19), further segregation takes place, with the big ones going to certain areas and the small ones to areas where they are much less likely to get coated with chocolate. This means that at the end of the process there is an even bigger range of sizes (see Project 2 in Chapter 12).

The shape is also important. Where possible, sharp edges should be avoided as these are difficult to coat and will often show through the chocolate. Slightly convex shapes are also better than concave ones. As shown in Figure 7.20, convex ones can only touch at a single point and so are easily separated by the tumbling action. The liquid chocolate, on the other hand, can get within a concave impression and form a sticky layer which will attach it to the next
particle. This will result in a lot of doubles or multiple pieces within the product.

The temperature of the centre is also important. Colder centres will help the chocolate to set more rapidly, but too cold a temperature will cause it to be uneven and to crack off at a later stage. In addition the size of the centre will change with temperature according to its coefficient of cubical (or volume) expansion ($\gamma$). This can be expressed by the equation:

$$V_t = V_0(1+\gamma t)$$  \hspace{2cm} (7.1)
where $V_t$ is the volume at temperature $t$ and $V_0$ is the volume at $0\, ^\circ C$.

The coefficient is therefore the increase in volume when 1 ml of a substance is raised in temperature by 1 $^\circ C$. This is different for different substances and tends to be much higher for most centres than for the surrounding chocolate. This means that after the centres have been coated, the products should be stored and sold at a relatively constant temperature, otherwise the centre will expand and crack the outer shell, which is unable to expand by the same amount. This is illustrated in Project 17 in Chapter 12.

Many centres need smoothing before being coated in chocolate, whereas others need a protective layer putting around them, so that the oil they contain does not migrate into the chocolate, causing it to bloom. Yet others are fragile and would break under the weight of the other centres falling onto them, or deform, e.g. soft raisins change their shape so that the chocolate cracks off. All of these need pre-treating, often with a sugar (sucrose and glucose) together with starches, gelatines or gum arabic.

The chocolate is then sprayed onto the centres as they tumble over each other in the pan or drum. This action smooths it out over the surfaces. The high shear and lower temperatures also crystallise the fat, and untempered chocolate can be used. The viscosity is important, in particular the yield value. If this is too high it will cause uneven coatings and even stick the centres to the pan walls. Too low a yield value chocolate will not stick to the surfaces and will leave areas of the centre uncoated.

The first few layers of chocolate must be put on very carefully. As soon as a uniform layer is seen on the centres, the temperature can be reduced to help the chocolate set. It is then warmed again and this procedure is repeated, often three to five times. After this, the chocolate additions and cooling can take place at the same time. The product actually warms up from the latent heat given out by the setting fat and from the friction between the falling pieces. Too cold air will, however, set the chocolate before it has been smoothed by the tumbling action and cause the product to be lumpy.

Many of these products have a very shiny appearance. It was noted earlier in this chapter that moulded products look much glossier than enrobed ones, due to the fat setting against a flat surface. Gloss is due to light being reflected, as in a mirror, and not
being absorbed by any surface scratches or blemishes. Some moulded products in fact looked relatively dull owing to the scuffing they receive during the wrapping procedures. This would happen for a lot of these products too, if it wasn’t for the protective coating that is used to polish or seal them. A very common coating is made from shellac, the refined resinous secretions from the lac insect. This is diluted with alcohol and can be sprayed onto the product in the pan, in the same way as the chocolate. This solution will, however, interact with the chocolate, giving a poor product unless an inner coating of sugar syrup, glucose and colloid (gum arabic, starch etc.) is used.

### 7.5.2 Sugar Panning

The sugar-panning process is used to coat the chocolate, or similar centres, with sugar. For chocolate panning, temperature control is used to harden the coating; for sugar coating, it is moisture reduction. The coatings are added in water-based solutions and dry air is blown over them so that the water evaporates, leaving very fine crystals.

The centres can be made by feeding tempered chocolate in the gap between two cooled rollers (see Figure 7.21). These rolls have matching indents, so that a thin web comes out of the bottom,
holding together all the centres. This web is removed and the centres fully solidified ready for panning.

The pan or drum is the same as for chocolate coating. The solution is then sprayed in. There are two types of coating, *i.e.* soft coating and hard coating. In the former (Figure 7.22) a liquid is applied and allowed to coat the surface before being dried out by adding an absorbing solid material, which is often sugar. This type of coating is often used for jelly-bean and sugar-egg types of product.

Hard coating (Figure 7.23), as is used in Smarties® and M&M’s®, has the sugar dissolved in the water. In order to minimise the drying time, the concentration is such that it is close to

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**Figure 7.22** Representation of the soft-coating process.

**Figure 7.23** Representation of the hard-coating process.
being a saturated solution at the temperature of use. The rate of drying is important in order to avoid cracking and also not to trap in moisture, which will later migrate to the surface and cause blemishes.

Coloured pigments can be added with the sugar and, in order to obtain a shiny surface, a wax outer coating may be put on the outside to give a smoother surface.

REFERENCES

CHAPTER 8
Analytical Techniques

A wide variety of analytical techniques is applied within the chocolate making industry. This chapter reviews some of the more widely used ones that are needed to monitor factors such as particle size, moisture, fat content, viscosity, flavour, texture and crystallisation. This is intended to give an insight into the science behind the techniques themselves, rather than to describe a particular instrument.

8.1 PARTICLE SIZE MEASUREMENT

Both the size of the biggest particles and the total (specific) surface area are important to chocolate manufacturers. Instruments like micrometers and optical microscopes can measure the diameter of the biggest particles, whereas sieving provides the weight of solid material above a certain size. None of these, however, can record the surface area of the smaller particles. Optical microscopes can measure particles as small as 5 microns, but, below this level, diffraction of the light makes readings very uncertain. (Diffraction is the bending of a wave motion around the edge of an obstacle – see Figure 8.2). Two types of instrument can be used to measure smaller particles; these are based on electrical conductivity and laser light scattering respectively.

Both methods rely on dispersing the non-fat solid material in a liquid before the measurement is carried out, so as to separate the individual particles. In chocolate these particles are so tightly packed that these systems would otherwise be unable to differentiate between the different particles. The dispersing liquid must, of course, have no effect on the particles themselves. This means that
they must not contain any water, which would otherwise dissolve the sugar. Very often oil, or an organic solvent is used. A few grams of chocolate are placed in the dispersing liquid. In order to separate the particles, some form of vigorous mixing is required, but this too must not be so vigorous that it breaks the particles. Placing the mixture in a low-power ultrasonic bath normally disperses the particles satisfactorily.

The electrical system operates by passing this dispersion through a narrow tube with electrodes on the side. The electric conductivity will be different when there is a particle present from that if there is only the liquid in the tube. In addition, this conductivity will depend on the particle's volume and electrical conductivity. If it is assumed that sugar, cocoa and milk solids all have a similar electrical conductivity, then by passing particles individually through the electrodes, it is possible to obtain a volume distribution. The mathematical treatment assumes that all the particles are spherical, in order to obtain an equivalent diameter. This is not exact, but the measurements obtained are sufficiently accurate to be very useful to the chocolate maker. Because the particle size distribution varies from submicron to about 70 microns, however, and because the electrode sensitivity depends on the diameter of the tube, it is often necessary to carry out the measurements with at least two sizes of tube. This makes the procedure relatively slow, and to a large extent this type of instrument has been superseded by those based on laser light scattering.

The sample preparation is the same as described above. The dispersion is this time circulated through a sample cell, which has a laser beam directed through it (see Figure 8.1). The laser beam is expanded to a diameter of between 5 and 20 mm and then shone through the suspension. In this instrument there is no attempt to measure individual particles, but only the dispersion pattern from all the dispersed particles within the beam. This pattern remains stable even though all the particles within the beam are in motion. The individual particles do, however, diffract the light. The smaller the aperture or particle compared with the wavelength of the light (for a helium neon laser it is 0.63 micron), the more it is diffracted (Figure 8.2). A combination of all the different scattering patterns is then imaged by a lens and monitored by a photodiode array. By changing the focal length of this lens, different particle size ranges can be monitored. Typical instruments measure 0.5 micron to
90 microns with one lens, but if partly manufactured chocolate is being monitored, *e.g.* from a two-roll refiner, the focal length can be increased to change this to, say, 1 micron to 175 microns.

The diffracted patterns are compared with those that would be expect assuming a combination of Fraunhofer diffraction for the bigger particles (≥ 40 microns) and Mie scattering for those of a size similar to the wavelength of the light. Particles must, of course, diffract/scatter the light individually, and not behave as agglomerates. When the dispersion becomes too concentrated, some of the smaller particles will coincide to give the signal of a larger one. This means that before taking the measurement the dispersion concentration must be adjusted to give the correct signal.

Once again it is assumed that the particles are spherical in order to calculate the volume of particles within certain diameter ranges, particle volume distribution curves and specific surface areas. A typical printout from such an instrument is given in Figure 8.3.
Although restricted to particles within the detection range, the normalised data is produced within a few minutes and this technique is therefore very useful for quality control.

8.2 MOISTURE DETERMINATION

Chocolate contains approximately 30% fat, but only about 1% moisture. Although moisture can migrate through chocolate, it is a very slow process, so removing the moisture from large pieces would take a very long time. In order to measure the moisture therefore, the chocolate can either be dispersed in another medium before drying, or the water determined by a chemical reaction which determines the presence of the H$_2$O molecule.

Traditionally moisture is determined by mixing a known amount of chocolate with pre-dried sand in a small metal dish, which is then weighed. The dishes are then heated in an oven for a set time.
and temperature. This may be 98°C for 12 hours or a lower temperature for a shorter time but under vacuum, before removing the dishes and reweighing them. Many laboratories have their own standard procedures, which are normally reproducible within that laboratory. Although the loss in weight does not give an absolute value for moisture, as not all the moisture is necessarily removed and a significant amount of other volatile material may have been, it does give a good indication of the “free” moisture within the chocolate. It is this “free” moisture that has the very large effect on the chocolate’s flow properties.

An alternative method uses the Karl Fischer reaction to determine the “free” and “bound” moisture within the chocolate. That is the moisture that is removed by oven drying together with any that is part of the water of crystallisation, e.g. in the lactose monohydrate from the milk. The latter does not play a significant role in effecting the chocolate flow properties and so does not really need to be measured. This technique therefore normally produces a higher measurement for milk chocolate than oven drying, but it does have the advantage of normally being more reproducible when automatic titration systems are used.

The Karl Fischer reaction depends on the action of sulphur dioxide, iodine and water.

\[
I_2 + SO_2 + H_2O = SO_3 + 2HI \tag{8.1}
\]

This reaction takes place very easily in a solvent composed of methanol and pyridine, according to the following equation.

\[
I_2 + SO_2 + H_2O + CH_3OH + 3py = 2pyH^+ + pyHSO_3OCH_3 \tag{8.2}
\]

This can take place in solvents other than methanol, but the pyridine must always be present. The overall reaction means that one molecule of water reacts with one molecule of iodine.

In the automatic system a pre-weighed amount of chocolate is dispersed in a mixture of formamide, chloroform and methanol in the reaction vessel. This vessel is then sealed to prevent further moisture coming in from the surrounding air. The mixture is then continuously monitored by two platinum electrodes that are placed in it. If any free iodine is present, this will depolarise the cathode and so the current will stop flowing. The potential difference across the electrodes is therefore used to control the titration. Initially the Karl Fischer reagent (which includes pyridine) is added slowly via a
peristaltic pump, which is able to record accurately the amount that passes through it. When the reaction is complete and no moisture remains, the current will stop flowing. The instrument can then calculate the percentage of water present, based on the sample weight and the amount of reagent used.

8.3 FAT CONTENT MEASUREMENT

Devices based on the reflection of near infrared radiation off the chocolate surface are able to give immediate readings concerning its fat and moisture content. This radiation is however affected by other factors such as particle size and chocolate composition, which means that instruments need to be re-calibrated for each product. The calibration for fat can be carried out using the traditional soxhlet method to determine its content accurately.

The soxhlet method involves using a solvent to dissolve the fat out of the chocolate. The solvent is then evaporated, leaving the fat, which can then be weighed. This analysis can be carried out in a glass system, such as is illustrated in Figure 8.4.

A weighed amount of chocolate is finely divided into small pieces before being wrapped in a filter paper, which is then put inside a permeable thimble container. This is then placed in the central part of the extraction system, the top part of which is cooled by a water jacket. The solvent, normally petroleum ether, is poured into the bottom flask where it is heated by an electric mantle. As it boils, the ether vapour rises through the side arm and condenses in the middle section. Here it runs down into the middle container where it comes in contact with the chocolate and dissolves the fat. The fat-containing solution passes through the filter paper and thimble and collects in the container until it reaches level A, corresponding to the top of the siphon system. It is then able to run back into the bottom flask. The solid material within the chocolate is retained within the filter paper and the fat collects inside the flask at the bottom – the mantle is hot enough to boil the ether, but not the fat.

Once all the fat in the chocolate has been removed, normally after about 12 hours, the top sections are separated from the flask. A new section, the petrol distillation unit, is placed on top, and will collect all the petroleum ether and not let it drip back. The fat is therefore left in the bottom of the flask and can be weighed. The whole procedure should take place within a fume cupboard, with
great care being taken to ensure that electrical connections and other objects present are unable to produce sparks and cause an explosion.

8.4 VISCOSITY DETERMINATION

8.4.1 Simple Factory Techniques

In Chapter 5 it was shown that liquid chocolate is a non-Newtonian fluid and so its viscosity cannot be described by a single number.

Figure 8.4 Diagram of soxhlet fat extraction apparatus.
In a simplified form it can be described as having a plastic viscosity and a yield value. Simple instruments do exist which will give single point measurements, two of which are the ball fall and the flow cup viscometers, shown in Figure 8.5 and used in Project 5 in Chapter 12.

In the ball fall viscometer there is a round weight attached to a wire or rod. On the latter are two marks. The ball is dropped into a container of liquid chocolate, normally controlled to be 40°C. Accurate temperature control is very important as a small difference in temperature has a large effect on chocolate flow properties. The time for the distance between the two marks to be covered in chocolate is noted and relates to the viscosity. Normally this measurement is carried out several times, but the first reading is discarded. If a bigger ball is used it will fall faster, so a series of measurements can be obtained by using several different sizes, and a flow curve is produced which is related to that chocolate. The rate of fall is however relatively slow for most chocolates, and the readings tend to relate to the yield value rather than the plastic viscosity.

In the flow cup viscometer, on the other hand, the chocolate is moving much faster, so its results relate more closely to the plastic viscosity. Here the chocolate and cup are warmed to the measurement temperature (usually 40°C). The cup is held above a balance on which is placed the receiving container. The plug is raised and the time recorded for a set amount of chocolate to run into this
container. The thinner the chocolate, the faster it will flow and the shorter this time will be.

### 8.4.2 The Standard Method

In order to measure the viscosity over a wide range of shear rates (degrees of flow or mixing) a concentric cylinder viscometer is used. This consists of a cup which contains the chocolate and a central bob. This bob normally has either a pointed bottom or has a hollow underneath, which is full of air when the viscometer is used (see Figure 8.6). The reason for this is that when the bob or cup rotates the chocolate will try to prevent it from doing so. This is the situation shown in Figure 5.2, with two parallel areas a fixed distance apart, moving at different speeds. The viscosity is directly related to the force the liquid develops to stop this movement. At the base the velocity is not uniform, being zero in the centre and maximum at the edge. This makes calculating the viscosity difficult. These two designs (Figure 8.6) have therefore been developed so that almost all the related force comes from the parallel area between the bob sides and the cup, and almost none from the base.

It is also important that the chocolate is uniformly treated within the gap. If the gap is wide and the bob turns, only the chocolate near the bob is going to be affected. The chocolate near the cup wall may even remain stationary (see Figure 8.7). To make matters worse, the viscosity of the chocolate varies depending on how fast it is moving. This means that even in a relatively narrow gap between the bob and cup there may be a range of viscosities across it. In

![Figure 8.6](image.png) Two different types of viscometer bob that are used to measure the viscosity of chocolate.
order to get a uniform flow across the gap it must be narrow and usually the ratio of the bob to the cup diameter must be $\geq 0.85$.

The chocolate must be free from fat crystals before the measurement is taken. To ensure this, the chocolate is heated to $50 \degree C$ and then cooled back down to just above $40 \degree C$ before pouring it into the pre-warmed cup. The bob is then put in and turned slowly so that the chocolate comes to a uniform temperature. The speed of rotation is then increased, with readings of the retarding force being taken at intervals. The speed of rotation is also noted, and can be converted into a shear rate (shear rate = relative velocity/distance between the bob and the cup). The viscometer is kept at the maximum stirring rate for a short period and then readings are taken at the same points as the speed is reduced.

In some viscometers the cup rather than the bob rotates, and the force on the bob is measured. In others, a specific force is applied to the bob and its speed of rotation is measured. For all these the apparent viscosity can be calculated from:

\[
\text{apparent viscosity} = \frac{\text{measured stress}}{\text{rate of shear}}.
\]

The shear stress or the apparent viscosity can be plotted against the shear rate (Figure 8.8). Using mathematical models, e.g. the Casson model, these plots can then be extrapolated to estimate what the stress would have been just as the chocolate started moving, i.e. how much force would be required to start it moving, which is the yield value. The apparent viscosity at the higher shear rates also corresponds closely with the plastic viscosity.

In order to get reproducible results, great care must be taken with sample preparation, temperature control and in ensuring that the viscometer is correctly calibrated.
8.5 FLAVOUR

Flavour and texture can only be determined using trained sensory panels. There is no correct flavour, as can be seen by the wide variety of “house” flavours, and the fact that the best-selling chocolate bars are very different in different countries, e.g. Cadbury in the UK and Hershey in the USA. There are, however, instrumental methods that help the manufacturer to determine whether the product is correct. Very often these relate to determining off-flavours. Chocolate has a mild flavour of its own, yet it very easily picks up other, often unpleasant, ones. For example, more and more chocolate is sold in petrol stations, yet the atmosphere in the forecourts often has a strong odour, which can be picked up by the chocolate. Likewise, great care has to be taken in ensuring the correct packaging material, as chocolate will quickly take up a cardboard flavour or one from printing inks, which will make it unacceptable to the consumer.

There are many techniques which can be used to detect off-flavours and odours or to characterise different cocoas or determine their degree of roast. Instruments such as the artificial nose (this contains a series of sensors whose electrical conductivity/resistance changes according to the amount of specific molecules that are present) have been suggested for the control of cocoa

Figure 8.8 Change in shear stress and apparent viscosity with shear rate as measured for a milk chocolate.
liquor flavour treatment and conching. It is currently used to ensure that fats are free from taints. In this chapter liquid chromatography is described, as this technique is able to be used for a wide variety of purposes.

Chromatography is an analytical technique in which different molecules are separated using the principle that their speed of migration through a medium depends on their sizes. Perhaps the best known example of this is the separation of food colours into several different coloured components by dissolving them in water and then letting the solution absorb through a filter paper or other media (see Project 13 in Chapter 12).

Many chromatographic techniques pass the sample through a medium where some of its components are either dissolved or absorbed, and then record the times at which different constituents come out of the medium. If the medium is packed in a broad column, there are many different paths the sample can take, so some may go straight through whereas other molecules can take much longer. This gives poor resolution, and often it is better to have long, thin columns.

In high-pressure liquid chromatography (HPLC), a solvent is continuously recirculated through tightly packed columns at pressures between 30 and 200 bars. Ultraviolet or visible photometers are often used to detect the components of the sample when they emerge. If these are fluorescent, then fluorescence detectors, which are more sensitive, can be used.

The columns can be packed with solids, gels or porous or particulate materials. The pore or particle size should be kept as small as possible to increase the column efficiency. The solvent may be aqueous or of a modified hydrocarbon, depending on the sample type. The analysis can take place within a few minutes.

HPLC has been used to detect the phenols associated with “smoky” flavour in cocoa. Ziegleder and Sandmeier\(^1\) investigated roasted cocoa using an ultraviolet detector. They found five peaks to be of particular interest:

1. pyrazine
2. 2,3-dimethylpyrazine
3. 2,5-dimethylpyrazine
4. 2,3,5-trimethylpyrazine
5. 2,3,5,6-tetramethylpyrazine.
Peak 5 is associated with raw cocoa, whereas peak 4 correlates with the time of roasting above 120°C. Peaks 2 and 3 increase after long roasting times at high temperatures and are an indication of over-roasting.

With the increased use of vegetable fats in chocolate (Chapter 6 and 10), HPLC has been seen as one of the possible approaches to determining their levels of use. It can also be used to detect adulteration of fats or oils.

8.6 TEXTURE MONITORING

As with flavour, sensory panels are the best way of assessing the texture of a product. Panels are, however, expensive to operate – tasters can evaluate only a few samples per day. They may also be unable to detect small changes. A manufacturer may make a change to ingredients or processing and want to know how it will affect the texture of the final product. In this case, analytical techniques are especially useful, as they can evaluate a large number of samples with a high degree of reproducibility.

For chocolate, snap and hardness are two of its most important characteristics. These can be analysed by texture measuring instruments, such as the one shown in Figure 8.9. These are able to drive probes, or blades etc. into a sample at a constant speed or force, whilst at the same time recording the resistive force or distance generated by the sample.

The relative snap of different chocolates can be measured by conducting a three-point bend test. Here bars of the test chocolates are moulded into bars of the same thickness. They are also stored for an extended period at the same temperature to ensure that this is uniform throughout the bars. Small changes in temperature can have a much larger effect on texture than small changes in composition. In addition, chocolate is a very poor conductor of heat, so it takes a long time for the centre of the bar to reach the same temperature as the surface.

The bars are then placed across two parallel supports. A probe, also parallel to these supports, is driven into the centre of each bar from above. Often the probe and support are made from rods with the same radius (see Figure 8.10). The force versus distance graph obtained from the instrument can be related to the snap (Figure 8.11). A chocolate with a good snap will have a steep curve
Figure 8.9  A TA texture-measuring instrument.

Figure 8.10  Three-point bend test on chocolate.
and a short recorded displacement as the bar will quickly break. A chocolate with a poor snap will tend to bend and so give a long, slow rise.

For hardness testing, a point or a ball is driven into a bar of chocolate placed on the base plate of the instrument. This can be driven in at a constant force and will be a force versus penetration trace, which will give a curve similar to the one for the three-point bend test. A hard chocolate will once again produce a very steep curve, whereas a soft chocolate will give much less resistance so the gradient will be less steep.

8.7 CRYSTALLISATION AMOUNT AND TYPE

8.7.1 Nuclear Magnetic Resonance

When molecules are in the liquid state they can move much more easily than when they are bound as part of a solid matrix. If a magnetic field is applied, many molecules try to line up with it, the rate at which they are able to do so relates to whether they are bound or free. This principle is used to measure the solid/liquid fat ratio in fat and chocolate by nuclear magnetic resonance (NMR). This is more accurate for fat, as the other solid components in chocolate, in particular naturally occurring metals such as copper, reduce the sensitivity.
In pulsed NMR, the sample is held in a magnetic field, which causes the protons in it to become polarised. A short, single radio frequency pulse is then applied, which causes them to rotate about the field. After the pulse, the molecules return to their original alignment over a period known as the relaxation time. This is much longer for the molecules which are bound within a solid structure. Different frequencies can be used depending on the types of molecule being investigated, which can be either water or fat based.

This type of information can be used to build up three-dimensional pictures of what is happening inside an object in a non-destructive manner. This technique is known as magnetic resonance imaging (MRI) and is the principle of the body scanner used for medical diagnostics. This has also been used to trace the migration of soft fats through chocolate, as the soft fat is much more liquid at room temperature. Figure 8.12 shows an image of a selection of confectionery products. The fat in the centres is much softer (more

![Figure 8.12](image_url) Confectionery items as seen by MRI. The softer fat appears lighter. (Guiheneuf et al.)
liquid) than that in the chocolate, so it shows up as being much lighter on the image.

8.7.2 Differential Scanning Calorimetry

The types of crystal present within the chocolate can be measured by X-ray, but this is expensive and the equipment is not compatible with most industrial food factories. The cooling-curve type of temper meter shown in Figure 7.2 indicates whether the chocolate can be solidified satisfactorily, but tends to be operator dependent and does not indicate the types of crystal present. The operator dependence has been overcome by more sophisticated temper meters, which operate on the same principle, but use electrical cooling to give a more uniform cooling rate, and use computers to analyse the cooling rates. The differential scanning calorimeter (DSC), can, however, indicate the relative amounts of the different crystal types that are present.

This instrument is based on the principle that when a substance melts (or solidifies) a large amount of energy is required – the latent heat – but the actual temperature of the sample will be almost the same. For solid chocolate this can be used in the following way. A small (about 2–10 mg) sample of chocolate is placed in a metal container. This is then heated to give a constant temperature rise of, say, 5.0 °C per minute. The amount of energy required to do this is compared with that required to uniformly heat a control sample, e.g. a metal such as an empty container, where no melting will occur. When a particular crystalline state starts to melt, more energy is required to maintain the temperature rise, so a peak starts to rise on the energy versus temperature graph, as illustrated in Figure 8.13. The peak reaches a maximum at the temperature at which the rate of melting is greatest. As is also shown in Figure 8.13, several different peaks can occur for the same sample showing that more than one crystal type is present. A single peak with a maximum about 34 °C will be obtained when the chocolate is well tempered in Form V.

Sometimes it is necessary to monitor the types of crystal present during the setting stage. In this case the remaining liquid fat can be rapidly crystallised by plunging the sample into liquid nitrogen. This fat will be in the unstable, lower melting-point forms and so can be distinguished from those crystals which have already set.
REFERENCES


Figure 8.13  DSC traces for different crystalline states in chocolate.
CHAPTER 9

Different Chocolate Products

Chocolate is sold in many forms, and with specific recipes being required for people with special dietary needs. In addition, there are particular formulations for ice-cream coating and with shape-retaining properties for sale in hot climates. At first sight many products often appear very simple, e.g. Aero® (Figure 9.1), but how do you put air bubbles inside a bar? Also how do you put a multicoloured centre inside a coating that is made of different types of chocolate and then manufacture the sweets on an industrial scale?

9.1 SPECIAL RECIPES

There are many types of chocolate available on the market in which the sugar (sucrose) has been replaced by sugar alcohols or polydextrose (see Chapter 2). This reduces the calories (on a weight for weight basis), and with certain sugar replacers, such as xylitol, can even have a beneficial effect on teeth. Although having a similar taste and texture to normal chocolate, most have a laxative effect, so limited amounts should be eaten. Different claims are made for these bars.

Low Calorie. This claim varies from country to country, but in the UK means that it has 30% less calories than normal chocolate. This is normally achieved by reducing some of the fat (which has a relatively high calorific content), together with using sugar alcohols and polydextrose.

No Added Sugar. Here the sucrose has been replaced by some other sweet ingredients.

Sugar Free. Because there is lactose in the milk, this too must be replaced as well as the sucrose. Special lactose-free “milk” powders are commercially available.
Reduced/Low Fat. The current EU nutrition and health claims regulations state that to make a reduced-fat claim the product must have 30% less fat than normal chocolate. The fat however provides the chocolate with its texture and melting property, so it is very difficult to remove this amount and yet retain good eating properties. There are numerous patents in this area, but so far no successful commercial products.

To make a low-fat claim a product must have less than 3 g fat per 100 g product. This is not possible using normal chocolate ingredients.

Suitable for Diabetics and Low Carbohydrate. Once again these often involve the use of sucrose replacers such as sugar alcohols and polydextrose.

High-Cocoa Content. These obviously contain a high level of cocoa nib. Often this is from a particular country. It should be remembered that the term “cocoa solids” includes cocoa butter. This means that many high cocoa chocolates also have a high fat content.

9.1.1 Ice-cream Coatings

Traditionally choc-ices were manufactured using a coating made from cocoa powder and non-cocoa vegetable fats. The coatings tended to crack easily and fall away from the centre. If normal tempered chocolate were used, it would be very hard and not have a pleasant melting sensation within the mouth. This is because it
has been stored at freezer temperature (about –18 °C), so all the fat is solid and takes a long time to melt. In order to overcome this, ice-cream chocolates are made softer by adding a lot more fat, normally more than 10%, but to be called “chocolate” only cocoa butter, milk fat or cocoa nut oil can be used. The milk and cocoa nut oils in particular make the chocolate much softer to eat and less likely to crack. In addition the coating chocolate is added to the frozen product in the untempered state. This means that it solidifies in Forms I or II (see Chapter 6). Here the crystals are less tightly packed than for tempered chocolate, so the product is softer. In addition, they need only to reach about 16 °C before they melt (not 32 °C as in normal tempered chocolate), so they do so much more quickly in the mouth – or on your clothes if you are unfortunate enough for a piece of chocolate to fall off the bar onto them.

9.2 SHAPE-RETAINING CHOCOLATE

During hot weather, chocolate often suffers from the following problems:

1. fat bloom
2. loss of shape
3. sticking to the wrapper
4. pieces of the product stick together – especially common with biscuits
5. make the hand messy.

There are several ways of reducing bloom formation due to heat, some of which were described in Chapter 6. The other factors can in fact be more unpleasant to the consumer and several methods have been developed to deal with them:

1. modifying the fat phase
2. coating the product with an edible transparent film
3. using water
4. building a skeleton of solid particles.

9.2.1 Modifying the Fat Phase

In Chapter 6 it was explained that cocoa butter from Malaysia melts at a temperature several degrees higher than that from Brazil.
So by sourcing the appropriate cocoa butter, it is possible to slightly improve a product’s shape retention. In addition, vegetable fats with a higher melting profile, known as cocoa butter improvers, can be added.

The improvement is small, however, as the fat must liquefy below mouth temperature for the chocolate to have good eating characteristics. Any fat remaining solid at this temperature will give the product a waxy texture.

9.2.2 Transparent Coatings

Here the product is coated with a film, which is often sugar or shellac based. It gives the product a shiny appearance and stops it melting in the hand. The coating can be applied by spraying or during panning. Because of the difficulty in getting a uniform coating, it tends to be applied to round/oval-shaped products, which can be tumbled in a revolving pan.

9.2.3 Water

Water thickens liquid chocolate, with the sugar particles sticking to one another and stopping it flowing properly. This means that high-moisture chocolate will keep its shape, but it is very difficult to mould or enrobe, so it is necessary to find processing ways of getting around this. One way is to add water in the form of fine droplets just before the chocolate is moulded. If this is done quickly before the water has the chance of attaching itself to the sugar, then the chocolate remains thin enough to process.

If a product has a moist centre (e.g. Mars Bar®) it will become much more heat resistant than one with a dry centre such as a KitKat®. The reason is that some of the moisture migrates from the centre through the chocolate, dissolving the surface of the sugar particles as it goes and sticking them to the neighbouring pieces. It is equally possible for the moisture to migrate from the outside into the chocolate and a Mars patent describes putting chocolate products in water-permeable wrappers and then storing them in high-humidity rooms.

A third method is to add the water in the form of a water-in-oil emulsion, i.e. water droplets coated by an emulsifier such as lecithin, all surrounded by a fat such as cocoa butter. Care must
be taken when adding the emulsion to the chocolate as vigorous stirring will break the emulsion and release the water (see Project 4 in Chapter 12). When carried out correctly, however, the water will slowly migrate throughout the product.

Although water will give the product good shape resistance above the melting point of cocoa butter, the texture is often more crumbly than normal chocolate and it often has a poor gloss.

9.2.4 Building a Framework of Solid Particles

In chocolate, fat is normally the continuous phase (see Figure 5.9). This means that it is possible to go from one side of the product to the other simply by going from one fat molecule to the next. Several processes produce a second path through the product by joining the sugar particles together, which in effect makes a “skeleton” framework. If the fat in this type of product is removed, for example by placing it in ether fumes, then a crystalline framework will remain (see Figure 9.2). This framework is easily broken in the mouth and so the texture is largely unchanged. The product will retain its shape even above 50 °C, when all the fat is liquid. The fat also tends to remain within the framework and will therefore cause less staining of the wrapper.

There are several ways of creating the supporting framework. One uses a small amount of glycerol, but, as with water addition, the process must be carried out quickly before the chocolate

Figure 9.2 Crystalline sugar framework in shape-retaining chocolate.
thickens. A Nestlé patent replaces some of the crystalline sugar by finely ground amorphous sugar. Particles of this material take in moisture and then transform it to the crystalline form. As they do so they eject the water molecules again, which sticks them to the neighbouring particles, before also enabling them to crystallise too. Over a period of weeks this chain reaction produces the full framework.

Hershey, the USA manufacturer, made a shape-retaining chocolate product called Desert Bar, which incorporated egg white. Although the bar was almost all chocolate, it could not be labelled as such, because egg white is not a permitted ingredient in chocolate. It was therefore called by another name and labelled as “containing chocolate”.

9.3 AIR BUBBLES IN CHOCOLATE

For a bubble to exist, the pressure inside it \( (p_2 \text{ in Figure 9.3}) \) must be in equilibrium with the pressure outside \( (p_1) \) and with that due to the surface tension of the chocolate \( (T) \). If the diameter is \( r \), it is possible to consider the equilibrium of the forces on one half of it (labelled B). The external force on B is \( \pi r^2 p_1 \), since \( \pi r^2 \) is the area of the circular face of B and pressure is force per unit area. Similarly, the internal pressure is \( \pi r^2 p_2 \). The surface tension can act only

![Figure 9.3](image-url)  
**Figure 9.3** Diagram illustrating pressure within a bubble inside chocolate.
around the circumference of the bubble, which has a length of $2\pi r$. It is then possible to find an equation that balances the forces:

$$2\pi T + \pi r^2 p_1 = \pi r^2 p_2$$

(9.1)

This simplifies as

$$p_2 - p_1 = 2T/r$$

(9.2)

The surface tension $T$ is a function of the type of chocolate and its viscosity, so this means that for a given chocolate the smaller bubbles will have a higher pressure inside them than the larger ones, for the same outside pressure. If two bubbles come together, then the pressure will move from the high-pressure bubble to the low-pressure one, with the result that the large ones will grow at the expense of the small ones.

Very importantly, if the difference in pressure ($p_2 - p_1$) increases, then so must the radius of the bubble in order to maintain the balance of eqn 9.2. This means that if you take a liquid chocolate under atmospheric pressure and then place it under vacuum, any small bubbles will grow dramatically, provided the chocolate is thin enough to move. In addition, more air will be released from the fat phase, creating new bubbles and enlarging others. This is because the solubility of a gas in fat increases with pressure. If the gas had been in the saturated condition at the original pressure, then the more the pressure is reduced, the more gas is released.

Changing the difference in pressure can be done in two ways. In one, the liquid chocolate is put in a mould and then placed in a cold vacuum box. The air bubbles form very rapidly and then the chocolate sets, preventing it from collapsing again. Sometimes additional gas, usually carbon dioxide, is mixed into the chocolate before the vacuum is applied.

In the second method, the gas is vigorously mixed under high pressure with the liquid chocolate. This dissolves the gas and forms micro-bubbles, both of which form the larger bubbles in the product when it is deposited into a chocolate shell and returns to normal atmospheric pressure. The chocolate is then cooled so that the bubbles are preserved.
9.3.1 Factors Affecting Bubble Size

Viscosity. Figure 9.4 shows some of the shapes and sizes of bubbles that can arise when changing the viscosity of the chocolate before placing it under a vacuum processing system. The change can be made by altering the emulsifier, fat content, moisture, or any of the other factors described in Chapter 5.

Pressure Difference. This is predicted by eqn 9.2.

Rate of Change of Pressure. This is particularly important in the vacuum process.

Rate of Set of the Chocolate. If the chocolate sets quickly the bubbles don’t have time to burst into each other and so there are far more of the smaller ones.

Type of Gas. Some gases like carbon dioxide and nitrous oxide are far more soluble in the fat than others such as nitrogen. When making aerated chocolate by mixing gas under high pressure the more soluble gases are required in order to give bigger bubbles. Work at Reading University also showed that the type of gas changed the taste, with a preference for the nitrous oxide-produced samples over the carbon dioxide ones.

When nitrogen is used the bubbles are very small and often not visible to the naked eye. This reduces the density of the chocolate, but not of course its calorific value (calories per gram). With a reduction in density of about 10% it is very difficult for the consumer to detect any difference between it and normal chocolate.

Figure 9.4 Different bubble sizes due to changes in chocolate viscosity.
chocolate. Reducing the density still further increases the choco-
late’s creaminess, makes it melt more quickly and it becomes lighter
in colour.

9.3.2 Water Evaporation Bubbles

Although not normally used commercially, it is possible to produce
bubbles in chocolate by adding water and then evaporating it, usu-
ally under vacuum. This can be done, for example, by emulsi-
fying the water with the chocolate and then spreading the emulsion
into sheets. These are then freeze-dried before being cut into pieces.
Figure 9.5 shows schematically the water-containing chocolate.
The holes vacated by the water droplets become the bubbles in the
product. This can be very light, almost half the density of tradi-
tional aerated chocolate. It melts very quickly and is often whitish
in colour.

![Figure 9.5](image)

Figure 9.5 Schematic diagram of water in cocoa butter emulsion in chocolate
matrix.
9.4 CREAM EGGS AND OTHER FILLED CHOCOLATE SHAPES

Chapter 7 described a standard moulding line to produce a chocolate shell, fill it with another ingredient such as caramel, pour on the chocolate back and then cool it. Figure 7.7 lists the 16 processes involved, which makes it a rather expensive and time-consuming procedure. For the mass production of products like cream eggs and praline-filled spheres (Figure 9.6) a much simpler process exists, which is limited to mould heating, depositing and vibrating, and cooling and de-moulding. This is done by a machine known as a single-shot depositor.

The principles of operation of this type of depositor are shown in Figure 9.7. There are two nozzles, an inner one, which deposits the central material and an outer one for the chocolate. The outer one starts first and enough chocolate for the base is extruded towards the mould. The inner nozzle then starts and deposits all the centre material. During this time the chocolate continues to come through the surrounding outer nozzle. The inner one then switches off leaving the top chocolate to be deposited by the surrounding ring. The whole product is then vibrated into shape. The deposit can be made directly into a tablet-shaped mould directly onto a belt, or into any shape of mould (see Figure 9.8). Once deposited, a cooling time of at least 40 minutes is required to ensure that the product doesn’t crack.

Figure 9.6 Typical product produced by single-shot depositor.
The timing of the pistons is very important, as is the positioning of the nozzle in the mould for some products. There are, however, several other critical aspects of this procedure, which limit the types of products that can be made by it.

Temperature of the Chocolate and Centre. The temperature of the centre must not be much higher than that of the enclosing tempered chocolate. If it is, the chocolate will de-temper, resulting in slow setting and poor contraction, making it hard to de-mould. The product will also have a poor texture and will easily bloom. The actual temperature that can be used will depend on the type of chocolate and its fat phase. This temperature is higher for dark chocolate at about 33–36 °C than for milk chocolate, which will be in the range 28–31 °C.
The Viscosity of the Chocolate and Centre. The process operates best when the viscosity of the centre material is similar to that of the surrounding chocolate. The latter can be increased by lowering the temperature and/or increasing the temper. Originally single-shot depositors were mechanically operated cam systems, which required very similar viscosities. More modern programmable electronically controlled machines enable products to be made with bigger differences in viscosity.

Nuts, Raisins and Other Pieces. The centre mass is forced through the inner nozzle. This means that it is possible to include only small pieces, and it cannot be used with whole nuts, cherries or raisins etc. Care must be taken with small pieces as these can block the central channel or break the outer chocolate surface.

The Ratio of Filling to Chocolate. Mechanical depositors limited the centre to about 40% of the product. Electronically controlled machines normally produce them with about 55% filling, although 80% may be achievable in certain circumstances. The shape is also important, with egg-shaped products having higher centre proportions than moulded tablet shapes (Figure 9.8).

Leaking Centres and Sticking of Sweets. Tailing occurs when the centre, instead of stopping flowing immediately the depositor piston stops, forms long “tails” or “strings” due to its elastic or gummy nature. These often extend into the chocolate (Figure 7.9) and may pass through it, eventually leaking out of the sweets and sticking them together. This is a particular problem with soft caramels, syrups and jellies.

Figure 9.9 Chocolate products with two components in the centre and shell (Awema A.G., Switzerland).
9.5 MULTIPLE CHOCOLATES AND CENTRES

The single-shot technology has continued to be developed over the years. By using a mechanical nozzle plate and the specially designed hopper dividers, the Swiss Awema company have made it possible to produce products with two fillings in two component shells, as shown in Figure 9.9. With this machine, the viscosity of the two fillings can be quite different.
CHAPTER 10

Legislation, Shelf Life and Packaging

Chocolate is controlled by very strict regulations in most countries. What is present is normally described on the label together with the nutritional information. Although confectionery products tend to have a relatively long shelf life compared with most foods, care must be taken with the recipes used for the centres in order to minimise changes caused by fat and moisture migration. The packaging itself will also play a major role in determining a product’s shelf life. It is also what a consumer first sees in a shop and must therefore be attractive as well as practical.

10.1 LEGISLATION

The actual legislation that is applied varies from country to country and is constantly changing as attempts are made to standardise it within trading areas, such as the EU. Most standards have limits as to the minimum amounts of cocoa and milk that must be present. These are referred to as milk and cocoa solids respectively. This can be a bit misleading as “milk solids” includes milk fat, which is a liquid at room temperature, and similarly “cocoa solids” includes any cocoa powder, cocoa liquor and cocoa butter. The legislation also stipulates the minimum amount of these fats that must be present and also the composition of the milk. It is not possible just to have lactose or whey instead of milk powder. The EU legislation also regards a higher milk (and therefore lower cocoa) content as being different and this has to be called “family milk chocolate” in all countries except the UK and the Irish Republic, where
traditionally it has been very common. This must still contain a minimum of 20% of both cocoa and milk solids, compared with minimums of 25% and 14% (cocoa and milk respectively) in normal chocolate.

If a chocolate contains vegetable fat, this must be made from only illipe, palm oil, sal, shea, kokum gurgi and mango kernels, used singly or in combination. These are all cocoa butter equivalents and could be used at higher levels without detriment to the product. The labelling would have to be changed however. They must also have been obtained by refinement and/or fractionation. Processes such as hydrogenation and enzyme inter-esterification are strictly forbidden. Any product containing up to 5% of these six vegetable fats must bear the statement “contains vegetable fat in addition to cocoa butter”. This statement has to be in the same field of vision as the list of ingredients, but separated from it and in lettering at least as large as the product name.

The USA legislation is very different. A product described as “sweet chocolate” corresponds to EU chocolate, but is defined as containing a minimum of 15% by weight of cocoa liquor (mass). Standards also exist for buttermilk chocolate, skim milk chocolate and mixed dairy product chocolates, which are not very common elsewhere.

A third set of standards is the Codex Alimentarius (Latin for “food code”), which was established by the Food and Agricultural Organisation of the United Nations. Although having a lot of similarities to EU legislation, differences are present and must be taken into account when exporting to countries using this standard, e.g. the majority of Latin American countries, other than Colombia.

In Europe the presence of lecithin and of other types of emulsifier must also be printed on the label. These may be recorded as an E number, e.g. E322 from soya is soya lecithin and E442 is YN, an ammonium phosphatide emulsifier. The E number shows that it has been tested and shown to be safe.

The possible presence of allergens must also be given on the label. These include egg, gluten and lactose, but perhaps the most widely known are nuts, in particular peanuts. If any manufacturing machinery has been used to process nuts, all future product must be labelled “may contain nuts” (even if no more have been added) until stringent cleaning has been carried out.
10.2 SHELF LIFE

The shelf life of a chocolate confectionery product is determined by when its taste, texture or appearance changes by an amount to make it unattractive to the consumer. In many cases this is when it turns white due to fat bloom. This change in appearance is often coupled with a hardening in texture and a slower melting, caused by changes in the fat crystallisation (usually the Form V to Form VI transition). These changes are slowed down by good storage conditions and the use of special fats and emulsifiers in the chocolate or soft-fat filling, as described in Chapter 6.

The surface can also look unattractive if it is badly cracked. This was noted to occur for panned products, when the centre expands more than the chocolate coating due to changes in temperature. This will also, of course, happen for moulded and enrobed products, and so a constant temperature is desirable. However, moisture transfer can also cause the centre to expand, especially for some baked centres.

Although a continuous chocolate coating slows down moisture transfer, water molecules will migrate slowly through it, so expansion of this type of centre will eventually occur. Once a crack has appeared, the moisture transfer increases rapidly and the product’s appearance will deteriorate rapidly, as will the eating properties of the centre.

Wafer-like centres take up moisture and become soggy, loosing their crispness and become unpalatable. With high-moisture centres such as caramel or fondants, drying out can occur making them hard or gritty, so once again moisture transfer is a problem. This can be made worse when both types of material are present in the centre and they don’t have a chocolate or fat layer between then to slow down the transfer. An example of this is Lion Bar\textsuperscript{\textregistered} with a wafer centre and a caramel coating.

Figure 10.1 illustrates some of the fat and moisture migrations that take place and which lead to the deterioration of the product quality. Even water going into the chocolate can change the texture and make it taste staler, although this is relatively unimportant compared with the wafer/caramel reaction. The driving force that governs the rate of these changes is the difference in equilibrium relative humidity (ERH) \(=100 \times \text{the water activity}\). This can be measured for each of the components by placing them individually in sealed containers with a small air space above them. This air will
rapidly take up or give out moisture to the ingredient so that the two rapidly reach an equilibrium. This is then measured by a probe calibrated to read relative humidity. The ERH will depend very strongly on the moisture level within the ingredient, its composition and also the temperature. For the product, the important temperature is that in which it is going to be stored, which might be 15 °C for the warehouse and 22 °C for the shop.

In order to minimise moisture transfer in products as illustrated in Figure 10.1, the ERH of each of the three components must be as close together as possible. The wafer can have only a limited amount of water present before the texture becomes too soft, and the caramel will be too hard if too little is present. This means that the components must be modified to change the ERH. Small amounts of ingredients which are highly hydrophilic (water loving) can, for instance, greatly reduce the ERH.

The form of the packaging is also important as, unless a good moisture barrier material is used, the humidity of the storage area can play a big role, with dry centres becoming soggy under high humidity, and moist ones becoming hard or gritty in dry conditions.

10.3 PACKAGING

It is very important for the packaging to be chosen to take into account the product and the conditions under which it is going to
be stored. At first sight it might seem best to package a high-moisture centre with a water impermeable barrier. This is not always the case, however. Some years ago a Turkish Delight product was wrapped using such a moisture barrier only for a lot of the product to go mouldy. The reason behind it turned out to be connected with the fact that the temperature varied during storage. Many shops are warmer during the day than at night. The moisture from the Turkish Delight eventually migrated through the chocolate so that the air within the wrapper was saturated during the day, when the room was hot. When the temperature fell below the dew point, water condensed onto the packaging and the chocolate, giving rise to sugar bloom and mould. A method of testing packaging materials for their moisture and odour-barrier properties is described in Project 14 in Chapter 12.

A high proportion of chocolate products are bought on impulse. That means that the customer does not arrive at the selling place with the intention of buying it, but does so because they see it displayed. This means that it is important to the manufacturer to have products that are packaged attractively and show up due to bright colours etc. In addition, the more space that is available, the greater the likelihood of the customer seeing the product. Shelf space is limited in supermarkets, so alternatives are used such as bags that hang on hooks or large containers that are placed at the ends of rows of shelves or near checkouts.

There are many types of packaging, some just providing protection against dirt, whilst others provide a good barrier against external odours. Others become associated with the product, e.g. the Smarties® tube and Toblerone® (Figure 10.2). When the original Smarties® tube with the plastic cap was replaced, the hexagonal tube was used as it had a similar appearance. In order to meet production requirements (over 17,000 Smarties® are eaten in the UK every minute), these tubes are filled and sold by weight. As not all Smarties® are the same size, there may be different numbers of sweets in different tubes, and it may be possible to have 10% extra weight, yet fewer Smarties®. In addition, not all the colours may be present, as each colour is made separately and then they are all mixed. This can give some interesting mathematical problems when special new Smarties® types are introduced (see Project 12 in Chapter 12).

In addition there is pressure to reduce the amount of packaging material used. This is partly for environmental reasons and also
because it increases transport costs, as well as, of course, to reduce the cost of the packaging material itself. Reducing the thickness of the material is one way of doing this. The size of the wrapper is also important. Because so many chocolates are sold, only a few millimetres less of packaging per sample can save a lot of wrapping material.

Packaging machines are complex and expensive. In the 1950s large amounts of labour were required to package products, but now individual machines may wrap several hundred bars per minute. These machines are expensive, however, and it is important that they are kept running as long as possible. This not only means feeding a continuous supply of the product of the correct size into the machine (too large pieces may block it), but also ensuring that stoppages do not occur when the wrapping material roll runs out (most modern machines go automatically onto a new reel), or if the quality of the wrapping material is not consistent, when it stretches or breaks.

10.3.1 Foil and Paper Wrap

The traditional packaging for moulded chocolate blocks and tablets is foil and paper (Figure 10.3). The aluminium foil provides some protection against dirt, insect infestation and taint, whilst the paper can be brightly coloured. The product name and the required legal and nutritional information are also printed on it. The
thickness and size of the foil and paper can be minimised, and both these materials are easy to recycle.

In order to transport them, generally two to six dozen bars are placed in a cardboard box, known as an outer. These outers are made of paperboard and can be decorated on the outside so that the retailer can sell directly from them (Figure 10.4). Where extra shelf life is required, particularly in hotter, more humid climates, this outer can have a barrier film put around the outside.

Outers are combined together in shipping cases and put on pallets. In some cases outers are palletised directly. The pallet load is secured by having plastic film stretched tightly around it.

Foil is also used to pack seasonal novelties such as Easter eggs and Father Christmases. For the latter, the packaging is more complicated as the printed design on the foil must fit with the markings on the chocolate.

10.3.2 Flow Wrap

A large proportion of chocolate confectionery is sold as a “count-line”. This is an individual unit often purchased and eaten by the

Figure 10.3 Traditional foil- and paper-wrapped chocolate products.
The consumer as a snack and not as part of a meal. The majority of these are packaged using flow wrap. This has the advantage that large numbers of items can be wrapped by a single machine, over 500 a minute in some cases. The pack is sealed tightly and, with the appropriate choice of material, can be a very good barrier to moisture and odours. A similar procedure (although not really fitting the flow-wrap definition) is also used to make larger bags that contain a selection of smaller sweets, which themselves may or may not be wrapped. Typical examples of flow wrap packaging are shown in Figure 10.5.

Wrapping material is supplied as long rolls of printed material. Great care must be taken with the type and application of the inks, otherwise they can impart an unpleasant flavour to the chocolate. Space is often left so that the “best before” date can be added on each individual item as it is packaged. This may be done by ink-jet coding or laser burning. A wide range of materials can be used, including thin co-extruded or white, cavity polypropylene and film/foil/ionomer laminates.
The flow wrap is initially unwound from the roll and made into a tube (see Figure 10.6) by sealing it with heat or pressure. The product is fed into the tube, which is then cut into the required lengths. The open ends are then sealed, once more by heat or pressure. Sealing by pressure requires cold adhesive to be applied to the reverse side of the wrapping material during the printing process.

This sealing is particularly important where it is necessary to have a good moisture or taint barrier. Where heat is used to do this, sufficient time must be allowed for the sealing layers to fuse together, which not only restricts the packaging rate, but also introduces the possibility that the product itself will be damaged by the heat. A method of cold sealing has therefore become
increasingly popular. This is based on natural rubber latex combined with resins. Not all packaging films will accept this cold seal, however, and a release lacquer or film may be required to avoid blocking on the reel. Fast packaging with very good barrier properties has been obtained with cold-seal and metallised film (film with a microscopical aluminium layer deposited under vacuum).

### 10.3.3 Biopolymers

Environmental issues with regard to packaging are becoming increasingly important, in particular with regard to their properties as renewable materials (regenerated within a human lifetime) and their biodegradability (can be assimilated by micro-organisms). In addition to not having these properties, conventional plastics are made from oil and gas feedstock, which are both finite resources. This has led to the commercialisation of a wide variety of renewable biopolymers for packaging.

There are four main categories:

- starch based
- polyhydroxy(alkanoates/buty rates)/polyesters
- polylactic acid (PLA)
- cellulose based.

PLA is a polylactic acid produced from cereal starch. Aliphatic polyesters are produced by Cargill from cornstarch to be used for
flexible films and rigid packaging. Properties of PLA include high stiffness, transparency and gloss, good folding properties and twistability, and low water-vapour barrier.

Plantic is also produced from cornstarch. It is modified by the inclusion of plasticisers and processing aids before being extruded. Currently it is being used for vacuum formed trays (VFTs) for chocolate assortment boxes. It is rigid, resistant to static electricity and dissolves completely in warm water. The latter means that storage conditions for the finished product are critical as it can pick up moisture and become soggy, or alternatively dry out and become brittle.

10.3.4 Robotic Packing

Placing individual sweets inside a selection box is very labour intensive. Although this is still done by hand in many factories, in some this is carried out by robots. Image recognition systems identify that the sweets are the correct shape and which way they are pointing. An arm, often with a suction system at the end, picks up the sweet and places it in the proper location in the box. Early robotic systems were very restricted in that they were difficult to reprogram when sweets were changed within the selection.

Figure 10.7 Robots packing boxes of chocolates. (Aasted.Mikroverk ApS Denmark).
Modern, more powerful computing systems have overcome this problem (Figure 10.7).

Much simpler robots are used to manipulate boxes or packages of sweets and put them into outer packages. This type of machinery has dramatically reduced the number of people working in the chocolate industry and, together with the new processes described earlier, have transformed it from a high manpower craft industry to a higher production highly scientific and technical one.

REFERENCE

CHAPTER 11

Nutrition and Health

Dedication. This chapter is dedicated to the memory of Dr Nicholas Jardine, who was passionate about the nutrition of chocolate.

Food is required to give us energy, and chocolate is relatively energy dense, meaning it is high in calories for a small portion size. Because of this, it has often been included in the food supplies for polar explorers and lifeboat rations etc. It also contains the three essential components of food, i.e. protein, carbohydrate and fat (although not in ideal proportions), together with some vitamins and several minerals. Eating a limited amount (up to a standard bar per day) of chocolate has been shown not to have a significant effect on migraines, acne or tooth decay. On the other hand, cocoa has been found to contain compounds that have positive effects in the prevention of heart disease and possibly some cancers, whilst the possible psychological effects have had a lot of media attention.

11.1 NUTRITION

The protein, carbohydrate and fat content varies with the chocolate type, as does the amount of micronutrients present, e.g. minerals and vitamins. A 100 g bar of plain chocolate is able to supply 24% of the copper needed in a healthy diet, whilst milk and white chocolate are a relatively good source of calcium, which is widely regarded as being beneficial in maintaining strong bones. Typical nutritional values of these components for the different types of chocolate are given in the Table 11.1.
It is important to have a balanced amount of the different nutrients, which means having a varied diet. Although eaten mainly as a snack or treat, confectionery (including sugar sweets) provides about 14% of the energy of the average UK diet (about half of this 14% is from chocolate). This can be broken down approximately as follows.

<table>
<thead>
<tr>
<th>Nutrient</th>
<th>Plain</th>
<th>Milk</th>
<th>White</th>
</tr>
</thead>
<tbody>
<tr>
<td>Energy (kcal)</td>
<td>530</td>
<td>518</td>
<td>553</td>
</tr>
<tr>
<td>Protein (g)</td>
<td>5</td>
<td>7</td>
<td>9</td>
</tr>
<tr>
<td>Carbohydrate (g)</td>
<td>55</td>
<td>57</td>
<td>58</td>
</tr>
<tr>
<td>Fat (g)</td>
<td>32</td>
<td>33</td>
<td>33</td>
</tr>
<tr>
<td>Calcium (mg)</td>
<td>32</td>
<td>224</td>
<td>272</td>
</tr>
<tr>
<td>Magnesium (mg)</td>
<td>90</td>
<td>59</td>
<td>27</td>
</tr>
<tr>
<td>Iron (mg)</td>
<td>3</td>
<td>2</td>
<td>0.2</td>
</tr>
</tbody>
</table>

It is important to have a balanced amount of the different nutrients, which means having a varied diet. Although eaten mainly as a snack or treat, confectionery (including sugar sweets) provides about 14% of the energy of the average UK diet (about half of this 14% is from chocolate). This can be broken down approximately as follows.

<table>
<thead>
<tr>
<th>Nutrient</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fat</td>
<td>15%</td>
</tr>
<tr>
<td>Protein</td>
<td>6%</td>
</tr>
<tr>
<td>Carbohydrate</td>
<td>17%</td>
</tr>
</tbody>
</table>

Children tend to eat more chocolate bars than adults, who on average consume more chocolate cakes and biscuits.

### 11.1.1 Fats

Fat is the most energy-rich, major component of chocolate, providing 9 kcal g$^{-1}$ compared with 5 kcal g$^{-1}$ for both carbohydrate and protein. Normally it makes up about 30% by weight of a chocolate recipe.

As well as providing energy, fats can also affect blood cholesterol levels in a way which may increase the risk of heart disease. Two types of cholesterol must be taken into account. A raised level of low-density lipoprotein (LDL) cholesterol (also known as “bad” cholesterol) and a reduced level of high-density lipoprotein (HDL) cholesterol (“good” cholesterol) indicate a higher risk of coronary heart disease (CHD), heart attack or stroke. These conditions occur due to the build up of cholesterol in the blood vessels, forming a hard, thick plaque (a condition known as atherosclerosis), making it difficult for blood to flow. In the arteries going to the heart, this restricts the blood flow and hence its oxygen supply,
sometimes resulting in chest pains. There is a greater chance of a blood clot forming on the surface of these arterial plaques. This could stop the oxygen supply to part of the heart, which would no longer function, resulting in a heart attack or, if it interrupts the blood flow to the brain, the result would be a stroke.

How blood cholesterol is affected depends on the type of dietary fat present. As was shown in Chapter 6, the fat in chocolate is mainly made up of cocoa butter and milk fat. Cocoa butter is composed of about 34% stearic acid which has little effect on cholesterol levels. About 34% is oleic acid, which is either neutral or lowers blood cholesterol, and a further 27% is palmitic acid, a saturated fatty acid which has moderate cholesterol-raising properties. Most of the remainder are polyunsaturated fats. The vegetable fats used in some chocolates are also stearic-rich. Those present in products labelled “chocolate flavoured coating”, or a similar phrase, may contain fats with very different compositions, however. Milk fat, on the other hand, with its greater amount of medium-length chain saturated fatty acids is not as beneficial as cocoa butter in this respect. Studies feeding people with a relatively high level of milk chocolate (280 g per day) showed no difference on cholesterol level compared with their normal diet.

In Chapter 6, the difference between cis and trans fats was explained. Trans fats are regarded as unhealthy in that they raise the LDL cholesterol and lower the HDL, thus having a doubly negative affect. Cocoa butter is virtually trans fatty acid free, although there is about 3–5% present naturally in milk fat, with the most being in the summer milk. Although good alternatives now exist, trans fatty acids are present in the cocoa butter replacers used in some countries as a cheap alternative to cocoa butter (these have been phased out by many international confectionery companies).

11.1.2 Carbohydrates

These are very important in that they make up about half the weight of normal chocolate. Most of this is sucrose, but there is also lactose in the milk powder, or it may be added as an ingredient on its own. Sometimes small amounts of glucose are present to reduce the sweetness, whilst fructose is used in some countries to make diabetic chocolates. Fructose is now being used to a lesser extent due to its potential negative health effects. Small quantities of other
carbohydrates are present as dietary fibre from the cocoa. The quantity and type of carbohydrate is important particularly for diabetics because of its effect on the glycaemic index.

The glycaemic index (GI) is a measure of the increase in blood glucose that occurs over the two hours after eating 50 g of carbohydrate. Glucose is absorbed rapidly after being swallowed, resulting in a rapid rise in blood glucose, and is normally the standard (GI 100) against which other foods are assessed. A long, high glucose reading can give rise to diabetes and heart disease. A lower GI food keeps this glucose level within a normal range. Table 11.2 gives the GI of some common foods. Starchy products, such as bread and potatoes, are easily digested producing a rapid rise in blood glucose. Since sucrose is 50% fructose, which has a low GI, it too has a relatively low GI. As fat in a food slows the absorption of carbohydrate, chocolate has an even lower GI. The high-energy density nature of chocolate may make it less attractive to people with a tendency for type 2 diabetes.

### 11.1.3 Proteins

Proteins are present both in the non-fat cocoa and milk particles. The milk protein has a much higher nutritional value than the cocoa protein as it has a greater proportion of essential fatty acids.

### Table 11.2 Glycaemic index of some common foods.\(^a\)

<table>
<thead>
<tr>
<th>Food</th>
<th>Glycaemic index (glucose=100)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose</td>
<td>99</td>
</tr>
<tr>
<td>Baked potato</td>
<td>85</td>
</tr>
<tr>
<td>Wholemeal bread</td>
<td>71</td>
</tr>
<tr>
<td>White bread</td>
<td>70</td>
</tr>
<tr>
<td>Sucrose</td>
<td>68</td>
</tr>
<tr>
<td>Raisins</td>
<td>64</td>
</tr>
<tr>
<td>Banana</td>
<td>52</td>
</tr>
<tr>
<td>Chocolate (all types)</td>
<td>43</td>
</tr>
<tr>
<td>Oranges</td>
<td>42</td>
</tr>
<tr>
<td>Fructose</td>
<td>19</td>
</tr>
<tr>
<td>Peanuts</td>
<td>14</td>
</tr>
</tbody>
</table>

11.2 OBESITY

Obesity is a major problem in many industrialised and developing countries. People whose body fat collects around the abdomen (apple-shaped people) are at a much higher risk of diabetes and heart disease than the general population; and even those whose fat accumulates on the limbs (pear-shaped people) can be at risk. Obesity can be defined by using the body mass index (BMI). This is defined as a person’s weight in kilograms divided by the square of their height in metres (kg m\(^{-2}\)). An individual can be considered obese if they have a BMI greater than 30, as shown in Table 11.3.

Although a tendency for obesity can be due to genetic factors or metabolic problems, generally obesity occurs when the energy intake from foods exceeds that being used to make the body function and in activity etc. This excess energy is then stored as body fat. The typical energy requirements for adults are shown in Table 11.4. These vary significantly according to age, activity level, build and other factors such as muscle mass.

Eating snacks is becoming an increasingly important part of the overall diet for many people and it is therefore interesting to compare the energy intake from chocolate with other snack foods (see Table 11.5). When deciding what to eat, however, other factors

### Table 11.3 Body mass index classification.

<table>
<thead>
<tr>
<th>Grade</th>
<th>BMI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Less than 20 – Underweight</td>
<td></td>
</tr>
<tr>
<td>Over 20 to 25 – Desirable or healthy range</td>
<td></td>
</tr>
<tr>
<td>Over 25–30 – Overweight</td>
<td></td>
</tr>
<tr>
<td>Over 30–40 – Obese</td>
<td></td>
</tr>
<tr>
<td>Over 40 – Morbidly or severely obese</td>
<td></td>
</tr>
</tbody>
</table>

### Table 11.4 Adult energy requirements (COMA\(^2\)).

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Female</td>
<td>1940 kcal day(^{-1}) Most occupations</td>
</tr>
<tr>
<td>Male</td>
<td>2550 kcal day(^{-1}) Moderately active</td>
</tr>
</tbody>
</table>
must be taken into account, e.g. a banana constitutes one portion of fruit – and fruit is an essential part of the diet.

Chocolate, however, represents approximately only 14% of the diet (in terms of the energy contribution), even in the UK, where consumption levels are high. Most dietary surveys have found no connection between obesity and chocolate eating. A UK national diet and nutrition survey of young people aged 4–18 years found that overweight children ate more food, but were no more likely to over-consume chocolate than any other food type. Lack of physical activity was a greater problem. Similar results were found in the USA.

A recent study at the University of Hull involved feeding a cohort of adults an extra 45 g of chocolate per day over a period of eight weeks. No significant increase in weight was noted. In addition, a Leatherhead Food International survey in the 1990s found a small tendency for high-chocolate consumers to have a lower BMI than non-chocolate eaters. This was believed to be due to the fact that such consumers had greater activity levels.

While most studies do not support a link between chocolate consumption and obesity, a recent report from the UK suggests that the consumption of high-fat sweet products as a whole may be a factor in obesity in some women. As chocolate is an energy-dense food which is high in fat and sugar/ its consumption must, of course, be limited.

### 11.3 TOOTH DECAY

Dental caries or tooth decay occurs due to acids destroying the tooth enamel and its underlying structure. This acid is formed by the breakdown of carbohydrates, such as sugar and starch, by
bacteria found in tooth plaque. Saliva is able to neutralise this and allow the teeth to start to repair themselves. The greater the number of times that you eat, the more acid is formed and the less the recovery time. The presence of fluoride seems to enable the teeth to withstand up to seven acid attacks per day.

Chocolate contains a significant amount of sugar and so would at first sight be thought to be very bad for the teeth, but compared with many other foods this is not the case. In fact it contains several components which reduce its cariogenicity.

11.3.1 Anti-Caries Factor in Cocoa

Tannins are compounds which contribute to the colour and flavour of chocolate and are part of a chemical group known as polyphenols. In the 1960s Stålfors found that defatted cocoa powder gave less tooth decay than whole cocoa powder and that removing tannins reduced its ability to inhibit decay. These tannins have been found to have an anti-bacterial and anti-enzymatic activity. This decreases plaque growth and inhibits acid formation.

11.3.2 Tooth-Friendly Milk Proteins

Plaque is formed when glycoproteins from saliva bind together with polymers secreted by bacteria. Glycoproteins from milk interfere with this binding process and so reduce the effect of caries-producing bacteria. This plaque reduction has the additional benefit in that it helps prevent gum disease.

Reynolds in Australia found that casein (one of the milk proteins) was also able to help remineralise the enamel after an acid attack. A possible explanation involves its ability to bind with calcium phosphate. Part of the casein molecule is hydrophilic (water liking) whereas other parts are hydrophobic (water hating). This allows it to bind to both the plaque and the bacteria. Once absorbed in this way, the casein breaks down to form amino acids, which resist further acid formation. Calcium also becomes available to remineralise the teeth. Casein can also bind to hydroxyapatite in the tooth enamel and remineralise it in this way.

Milk also has a buffering effect, *i.e.* it prevents the lowering of the acidity (pH). The lower the pH and the longer that it remains low, the more damage is caused to the teeth.
11.3.3 Oxalic Acid
Jackson and Duke\(^3\) showed that concentrations of 0.2–0.4% oxalic acid reduced further acid production by between 35% and 60%, by inhibiting enzyme activity. This bacterial enzyme converts carbohydrates into lactic acid, which causes tooth decay. Oxalic acid is present in chocolate at less than 0.1%, however, so its effect may be limited.

11.3.4 Oral Clearance
Chocolate differs from many foods in that it becomes liquid in the mouth due to the fat melting and the sugar dissolving. This enables it to be swallowed very quickly, provided there is no caramel or other sticky inclusions, leaving little sugary material for the bacteria to convert into acid. Nuts as inclusions reduce the clearance time of chocolate further, thereby reducing its cariogenic potential.

11.4 OTHER ALLEGED NEGATIVE REACTIONS
11.4.1 Migraine and Headaches
Migraines are much more severe than normal headaches and are often accompanied by flashing lights and preceded by mood swings, food craving or tiredness. Amines such as tyramine, histamine and phenylethylamine, which are linked with headaches, are found in cheese, peanuts, red wine and chocolate. Although many people blame chocolate for their migraine, several studies indicate that this is probably not the case. Moffat, Swash and Scott\(^4\) studied 80 subjects and found that only 13 were affected by chocolate alone. Only two of these were found to consistently respond to eating chocolate. A much more likely explanation is that the food craving due to stress, experienced before an attack, encourages the eating of foods like chocolate.

11.4.2 Acne
Acne produces black, yellow and red spots particularly on the face, back and chest and is due to a blockage of the flow of oil from the sebaceous follicles on the skin. The cause isn’t clear, but could be due to obstruction of the canals, excessive oil flow, bacteria and/or
hormones. Chocolate is often blamed, although doctors normally agree that chocolate and fatty foods are not a cause. A University of Missouri study in 1971 of acne suffers found 10% blamed chocolate. Study subjects were then given the equivalent of 230 g (about four bars) of chocolate a day for a week, but no increase in acne was found.

11.4.3 Allergies

An allergy is an altered or abnormal tissue reaction following an exposure to a foreign substance or allergen, normally a protein. Very few people are allergic to the proteins in cocoa, but milk, nuts and peanuts are much more likely to cause reactions. Some people are also intolerant to the lactose, which is found in milk, but may also be added as a separate ingredient. UK legislation requires the manufacturer to clearly label any product to show possible allergens. Only very minute traces of some nuts can have a dramatic effect on susceptible individuals, so if machinery has previously been used for a nut-containing product, all future confectionery made on it must be labelled as possibly containing nuts, even though none have been added during manufacture. Very stringent cleaning procedures must be carried out before this warning can be removed.

11.5 POSITIVE HEALTH EFFECTS

The Aztecs and Mayans believed that their cocoa drink had a positive health benefit and there have been many documented uses of cocoa as a medicine over the centuries. Recently there has been renewed scientific interest into the benefits of eating cocoa and chocolate, particularly in the respect of reducing the risk of cardiovascular disease. The effect of the fat in chocolate on cardiovascular disease has already been described, but these new studies are related to physiologically active plant compounds known as polyphenols and more specifically to a class of polyphenols known as flavanols.

Flavanols are known to have antioxidant and other potential health benefits. Cocoa contains the flavanols epicatechin and catechin as well as procyanadins, all of which appear to be able to be absorbed by the body in proportion to the amounts that they are eaten. Procyanadins are normally broken down in the gut to form
epicatechin before being absorbed. Polyphenols are found in a wide variety of foods such as fruit, vegetables, tea, coffee and red wine. Dark chocolate is shown to have more polyphenols than milk chocolate, because it contains more cocoa (one study has shown that polyphenols are less bioavailable in the presence of milk). High-cocoa content, dark chocolate would be expected to contain an even higher level. This is not always the case, however, as the amount of polyphenol present depends also on the type of cocoa being used and how it has been processed. Cocoa fermentation in particular can destroy a lot of the flavanols.

The cocoa flavanols have been shown to be able to limit the progression of cardiovascular disease by exerting anti-platelet (anti-blood clotting), anti-inflammatory and antioxidant activity. Platelet reactivity and aggregation-enhanced blood clotting can cause thrombosis. Flavanols can affect the reactivity and reduce the agglomerational tendencies of the platelets. At normal consumption levels, flavanol-rich chocolate has a similar effect to that associated with taking a low dose of baby aspirin.

Flavanols have been shown to suppress some inflammatory reagents and increase the anti-inflammatory nitric oxide. Nitric oxide is an important factor as it relaxes blood vessels thus ensuring a healthy blood flow. Endothelial cells line the entire circulatory system, and when these malfunction (due to risk factors of heart disease, e.g. smoking) the level of nitric oxide drops, indicating restricted blood flow. Studies have demonstrated an improvement in blood pressure as well as an improvement in blood flow following the consumption of flavanols from cocoa and chocolate due to their affects on nitric oxide.

In addition to being an indicator of the risk of coronary heart disease, LDL cholesterol can oxidise and injure the arterial walls. Eventually this may lead to plaque formation and atherosclerosis. Consuming cocoa and dark chocolate has been shown to reduce the risk of LDL cholesterol oxidation.

Flavanols also have antioxidant activity which is able to help the body counteract damage caused by free radicals, which eventually lead to life-threatening diseases such as cancer. The antioxidant activity from consuming flavanol-rich dark chocolate is comparable to or greater than that from other antioxidant-rich foods and drink such as green tea, red wine, blueberry and garlic (see Table 11.6 for some typical figures).
Although the cause is not known, the consumption of high levels of cocoa solids can have a beneficial effect on treating other diseases. In a 2005 study at the University of Hull, sufferers with ME were fed with a chocolate containing a very high level of non-fat cocoa, or with a bar with the same colour and number of calories, but containing only cocoa butter and none of the brown cocoa particles, as a control in a double blind trial. Whilst eating the high levels of cocoa, patients showed a significant improvement, some of whom managed to return to work, whereas those consuming the control remained unchanged.

11.6 PSYCHOACTIVE COMPOUNDS

Methylxanthines are a group of chemicals that are found in more than 60 plant species that are thought to be capable of having physiological effects. Only two are found in chocolate in any significant amount, i.e. caffeine and theobromine. Caffeine stimulates the central nervous system, but theobromine is only about one tenth as effective. The amount of caffeine present is, however, lower than from many other sources (see Table 11.7). Although
more theobromine is present, its pharmacological activity is much weaker than caffeine. It has, however, been used in conjunction with the drug phenobarbitone to treat nervous excitement and insomnia. It is also a stronger diuretic (increases water excretion through the kidneys), although this would not occur from the amount of theobromine present in chocolate.

Cocoa also contains many other physiologically active compounds including phenylethylamine (an amphetamine-like compound), anandamide (a cannabinoid) and tryptophan, which have all been thought to induce cravings for chocolate. However, for different reasons, it is extremely unlikely that these compounds could exert a mood altering affect. For example, both phenylethylamine and anandamide are present in chocolate in only very small amounts, so vast quantities of chocolate would need to be consumed to obtain the active doses. Tryptophan is present in higher quantities in other foods that are not craved. Further, only a very low-protein, high-carbohydrate meal will allow tryptophan to influence the brain, and the level of protein in chocolate is too high for tryptophan to have an affect.

There is, however, some evidence that eating chocolate can improve your mood. An article in *Food Manufacturer* in 2002 reported a study involving 1000 students at the University of Bath performed by the chocolate manufacturer Cadbury. The study found that 70% of the students who ate their chocolate described themselves as being happy, compared with 41% of the students who hadn’t eaten any chocolate at all. This is probably because very pleasant palatable foods trigger the release of brain endorphins which are feel-good chemicals.

Most people like chocolate, but is it possible to become addicted to it? From the chemical composition this is extremely unlikely. But
the combination of pleasant taste, melt-in-the-mouth sensation and smoothness make it extremely attractive and are good enough reasons in themselves to make people crave for it.

REFERENCES

CHAPTER 12

Experiments with Chocolate and Chocolate Products

The aim of this chapter is to outline a series of projects which can be carried out to demonstrate chemical, physical or mathematical principles. They have been designed to use only relatively simple apparatus and should be able to be adapted for students over a wide range of ages. Appropriate hazard analyses and safety precautions must always be applied according to the circumstances where they are performed.

PROJECT 1: AMORPHOUS AND CRYSTALLINE SUGAR

Apparatus:

Beaker.
Magnetic stirrer.
Thermometer sensitive to better than 0.1 °C.
Balance capable of reading to at least 1 g.
Granulated sugar.
Skim milk powder.
Boiled sweets, e.g. Fox’s Glacier Mints® (NB not pressed or tableted sweets such as Polo®’s which are more crystalline).

Aim:

To show how amorphous and crystalline sugar differ when they dissolve in water.
The crystalline sugar causes the water to cool down as energy is required to separate the molecules (heat of solution). Amorphous sugar is in an unstable state, however, and gives out energy when it changes to its stable, lower-energy crystalline state. This means that there is spare energy so the water becomes warmer.

**Procedure:**

1. Pour 10 ml water into a beaker and place on the magnetic stirrer.
2. Place the thermometer in the water and continue the stirring until the temperature is constant.
3. Break the boiled sweets into small pieces. (This can be done by placing them within a material bag and crushing them with a hammer. CARE: TAKE APPROPRIATE PRECAUTIONS.)
4. Weigh out about 10 g granulated sugar and also of the crushed amorphous material.
5. Drop this quantity of granulated sugar into the water and record the temperature for the next five minutes.
6. Repeat the procedure using the amorphous sugar. This time the temperature should rise.
7. The experiment can also be tried with skim milk powder. This contains lactose, which is normally in an amorphous state owing to the rapid spray-drying process. This normally gives a much greater temperature rise than the boiled sweets.

**PROJECT 2: PARTICLE SEPARATION**

**Apparatus:**

Several sheets of paper.
Scissors.
A small, narrow glass jar with a lid.
A stopwatch.
Dried peas, rice, lentils and sunflower seeds.

**Aim:**

To investigate some of the principles that are used to separate cocoa beans from stones and cocoa nib from shell and that are
the cause of the segregation of some centres during chocolate panning.

**Procedure:**

*Separation by Vibration*

1. Partly fill the narrow jar with dried peas.
2. Pour some of the rice so that it forms a layer on top.
3. Stir so that the two are well mixed and put the top on the jar.
4. Shake gently with a slightly swirling action. The peas should come to the top and most of the rice should fall to the bottom as shown in Figure 12.1.
5. Turn the jar over and shake again. The peas should once more be at the top.
6. Repeat with other materials such as lentils and sunflower seeds. Shaking the jar on its side can sometimes be more effective.

*Separation by Falling Speed*

1. Cut one of the pieces of paper in half. Then cut one of the halves into strips about 1 cm in width.
2. Screw another piece of paper into a ball.

![Peas and rice mixture in a small jar after shaking.](image-url)
3. Hold the uncut half sheet of paper horizontally at a height of at least 2 m. Let go and time it as it falls to the ground. Repeat this several times and determine the average falling time.
4. Repeat this with the screwed up ball. This will fall much faster, and shows why the plate-like shell of cocoa beans can be separated by falling speed – or upward suction – from the spherical nib.
5. Repeat the experiment with the paper strips. These will fall faster than the uncut sheet, but slower than the ball. The falling speed of a fibre is governed by its diameter and not its length. So, providing that they are long enough to be fibres (the length should be at least 10 times the diameter), all the thin strips of paper with the same width should fall at about the same rate.

**PROJECT 3: FAT MIGRATION**

**Apparatus:**

2 desiccators or sealed containers.
Printing/photocopier paper.
Silica gel.
Cocoa butter as obtained from Project 4. (Other hard fats should give the same results.)
Confectionery wrapping material.

**Aim:**

To show that moisture affects the release of cocoa butter from the cells within the cocoa nib.

**Procedure:**

1. Put silica gel in the base of one desiccator. Leave a piece of paper in a relatively moist environment.
2. Put a piece of paper into the desiccator and leave for several hours.
3. Remove the pieces of papers and put a drop of fat on each. The fat will remain as a globule on the wet paper, but spread out and sink into the dry one.
4. Repeat with the PVC-type film used to wrap a lot of confectionery. This will not let the fat sink into it under either condition and is why it has replaced cellulose as a packaging material for a lot of confectionery.

**PROJECT 4: COCOA BUTTER SEPARATION**

**Apparatus:**
- 2 × 1 litre beakers.
- Oven.
- Knife.
- Bars of chocolate.
- Stirrer.

**Aim:**
To separate cocoa butter from the chocolate and to show the effect of lecithin as an emulsifier in a fat-in-water system.

**Procedure:**
1. Use the knife to scrape the chocolate into small flakes.
2. Three-quarters fill one beaker with water at around 60 °C. CARE: THIS IS HOT.
3. Slowly pour the flakes into the water and let them settle on the bottom.
4. Gently agitate the flakes using the stirrer – DO NOT MIX VIGOROUSLY.
5. Place the beaker in an oven at 50–60 °C for 12 hours. CAUTION: IT IS HOT.
6. Remove the beaker from oven and let it come to room temperature. You should find that a yellow fat layer has formed at the top, which is a mixture of cocoa butter and milk fat.
7. Repeat the above steps, but stir very vigorously when the flakes are in the beaker. CARE: THIS IS HOT.

The lecithin in chocolate is attached to the sugar. When the sugar dissolves it can go into the water. The fat is the continuous phase in chocolate, and if it remains in large globules when it melts, then the
layer of fat will form. If it is stirred vigorously, the fat forms small droplets which can be coated with lecithin. These remain suspended within the water as an emulsion.

**PROJECT 5: CHOCOLATE VISCOSITY**

**Apparatus:**

Temperature controlled cabinet (40 °C).
Thermometer accurate to 0.5 °C.
Tall beaker.
Ball fall viscometer (a design for its construction is given in Figure 12.2).
Stopwatch.
Plastic funnel.
Balance capable of reading to 1 g.
Bars of chocolate.
Sunflower oil.

**Aim:**

To show the effect on the flow properties of chocolate of fat and moisture additions.

**Procedure:**

*Effect of Fat*

1. Melt the chocolate and keep it, the sunflower oil and the viscometer, plus any glassware in the cabinet at 40 °C and

![Figure 12.2 Schematic diagram of ball fall viscometer.](image)
allow it all to come to temperature. Check the temperature using the thermometer.
2. Fill the tall beaker almost to the top.
3. Use the stopwatch to record the time it takes for the ball of the ball fall viscometer to sink between the two markings on the bar or wire.
4. Repeat this four or five times, disregarding the first reading.
5. Repeat with different sizes of ball, if available.
6. Place the funnel on a stand so that any liquid in it will flow into a container on the balance (Figure 12.3).
7. With a finger or card under the outlet, partly fill the funnel.
8. Measure the time it takes for about half the chocolate to flow out.
9. Refill and repeat the procedure several times.
10. Add sunflower oil to the chocolate so that it makes up about 3% of the mixture. Stir very thoroughly so that the mixture is uniform (a food processor can be used if available) and repeat both types of measurement.
11. This can then be repeated with other levels of addition.

Figure 12.3  Set up for flow cup viscometer.
What effect does the extra fat have? Does it affect the ball fall measurement (predominantly yield value) as much as the flow cup (mainly plastic viscosity) measurement?

**Effect of Moisture**

12. Repeat the procedure given above for the sunflower oil, but this time use water. How do these results compare with the previous ones. See also Project 8.

**PROJECT 6: PARTICLE SIZE OF CHOCOLATE**

**Apparatus:**

- Microscope (if possible with polaroids).
- Sieve (less than 50 microns mesh).
- Balance capable of reading to 0.1 g.
- Micrometer capable of measuring 40 microns.
- Sunflower oil.
- Several different brands of chocolate.

**Aim:**

To investigate the largest particles within chocolate and see whether particle size affects its texture/flavour.

**Procedure:**

**Microscopy**

1. Place a small amount of chocolate on a microscope slide and disperse the solid particles in a transparent liquid that will not dissolve them, e.g. sunflower oil.
2. Put a cover glass on top of the dispersion on the slide and press gently to remove the air. (Pressing too hard will move all the particles to the edge.)
3. Calibrate the microscope to be able to determine the size of particles greater than 20 microns.
4. View the sample and determine the size of the biggest particles.
5. If the microscope is a polaroid one, cross the polaroids so that the field of view is dark when no sample is present. Re-examine the sample. Crystalline sugar is birefringent, which means that
it will show up as a bright image (Figure 2.10), whereas the cocoa and milk particles will remain dark. It is therefore possible to see whether the biggest particles are sugar or not.

6. Repeat for all the different chocolates.

**Sieving**

1. Disperse about 10 g of liquid chocolate in about 100 ml warm sunflower oil. (If an ultrasonic bath is available, this will help to disperse the particles.)
2. Pour the suspension through the sieve. This will retain the biggest particles.
3. The biggest particles can be defatted and weighed, or viewed on the microscope as described above.

**Micrometer**

1. Prepare a concentrated suspension of chocolate in sunflower oil.
2. Place a drop of the suspension on one of the jaws of the micrometer.
3. Screw the jaws of the micrometer together and take a reading. This should be done several times and the reproducibility of the technique determined, as it is very easy to screw too hard and break the particles.

**Flavour**

1. Get at least five people to evaluate the samples. Each sample should be given a value between 1 and 10 for grittiness, for creaminess and for cocoa flavour.
2. Determine whether there is a correlation between any of the size measurements and these three sensory parameters.

**PROJECT 7: EFFECT OF LECITHIN**

**Apparatus:**

Food processor or mixer.
Icing sugar.
Sunflower oil.
Lecithin (obtainable from many health food shops).

**Aim:**

To show the large effect on viscosity of lecithin in a sugar–fat mixture.

**Procedure:**

1. Mix 5 parts of sunflower oil with 2 parts of icing sugar in a food processor for five minutes.
2. Check the viscosity by estimating how hard it is to stir, or use the ball fall viscometer (Project 5) if then mixture is thin enough.
3. Add 5% by weight of lecithin (warm slightly if it is very thick) and mix in the processor for a further two minutes.
4. Remeasure the viscosity.

**PROJECT 8: CHANGING THE CONTINUOUS PHASE**

**Apparatus:**

Oven set at 40 °C.
Stirrer (ideally connected to a torque or a power meter, but stirring by hand and estimating how hard it is to mix is possible.)
Bars of chocolate.

**Aim:**

To show that small additions of water to chocolate make it thick by “sticking” the sugar together (Project 5), but larger additions make it thinner again as the water is able to form a continuous phase.

**Procedure:**

1. Melt the chocolate for at least three hours in an oven at 40 °C.
2. Stir, and estimate or record how hard it is to do so.
3. Add 2% by weight of water, mix thoroughly and then determine how hard it is to stir it.
4. Repeat the procedure until 30% water is present.
5. Plot the “stirability” against the water content.
PROJECT 9: CHOCOLATE TEMPER

Apparatus:

Thermometer that includes the range 20–32 °C and is accurate to better than 0.5 °C. An electrical thermometer is preferred as it will have a much smaller heat capacity than a glass one. Less accurate thermometer with a range including 50 °C 2 beakers and stirrers. Test tube with stopper and stirrer. Stopwatch. Hot plate. Bars of chocolate.

Aim:

To show that pre-crystallised chocolate sets differently from untempered chocolate.

Procedure:

1. Melt about 30 g chocolate and heat in a beaker to 50 °C. (CARE: THIS IS HOT.) Stir occasionally and maintain this temperature for about 30 minutes, to ensure that no fat crystals remain.
2. Pour about 10 g into the bottom of a test tube, and put the stopper containing the thermometer and stirrer into the top as shown in Figure 12.4.
3. Put the test tube in a beaker of cold water and plot the change in temperature with time once the chocolate is below 32 °C. Stir intermittently as the chocolate sets.
4. Cut or grate about 5 g chocolate into a powder.
5. Repeat the above procedure (steps 1, 2 and 3) until the chocolate is about 35 °C.
6. Add about 3 g of the powdered chocolate and stir vigorously. Start measuring the rate of cooling. (The powdered chocolate should “seed” the chocolate with Form V crystals, which should then make it set and make the temperature rise much more rapidly than was the case with the first sample.)
7. The experiment can then be repeated using iced water or with different amounts of chocolate.
PROJECT 10: HARDNESS MEASUREMENT

Apparatus:

Temperature-controlled cabinet.
Set of half kilogram and kilogram weights.
Retort stand.
Counter sink, i.e. a broad metal rod with a conical tip at one end.
Travelling microscope (although it should be possible to do the experiment with a ruler and a hand magnifier).
Bars of chocolate.

Aim:

To demonstrate that small changes in temperature can have a large effect on the hardness of chocolate.

Procedure:

1. Store some bars of chocolate at different temperatures for at least 12 hours. These could include in a refrigerator, in a warm room, and in temperature-controlled cabinets at 24°C and 28 °C.
2. Set up the retort stand so that it will hold the rod of the counter sink (through a loose-fitting tube) vertically above the sample being tested (see Figure 12.5).
3. Take one of the samples and place it gently below the point of the rod. The samples should be removed from the storage conditions just before testing so that the fat doesn’t melt or harden.
4. Carefully place a weight on the top of the rod and balance it there for a few seconds.
5. Remove the weight and lift the point of the rod out of the chocolate.
6. Move the sample slightly to one side and put the point of the rod on its surface.
7. Carefully place a bigger weight on top of the rod.
8. This procedure should be repeated several times, a note being made as to which mark corresponds to which weight.
9. Measure the diameter of each of the marks using a travelling microscope.

Relate the diameter readings to the weight applied and also to the temperature of storage.

Figure 12.5 Apparatus for determining the relative hardness of chocolate.
PROJECT 11: CHOCOLATE COMPOSITION AND PRODUCT WEIGHT CONTROL

Apparatus:

Balance capable of weighing to at least 0.1 g.
Bars of solid chocolate and countlines.

Aim:

To look at the chocolate composition of products on the market.
To investigate the weight of the bars compared with their declared weight and to see if there is any difference between enrobed and moulded products.

Procedure:

1. Survey the chocolate bars being sold in the shops with respect to their cocoa and milk solids.
2. Note any that indicate the country of origin of the cocoa. These are most likely to be found on plain chocolate, especially organic products.
3. Buy at least ten samples of individual products, as far as possible from different shops. (Samples from the same shop are likely to have been made at the same time, so differences will be less.) The samples should include moulded chocolates like solid bars and countlines such as Mars Bar®, Drifter®, Crunchie® and Lion Bar®.
4. Weigh all the samples and plot a graph of the number of bars within set weight ranges.
5. Calculate the mean weight, the standard deviation and the coefficient of variation (standard deviation/mean) for each product and compare this with the declared weight. The coefficient of variation of the moulded solid bars would be expected to be smaller than that for enrobed ones with centres, as not only is enrobing normally less precise, but there is also often a weight variation between the centres.
PROJECT 12: DISTRIBUTIONS AND PROBABILITIES

Apparatus:

At least 20 tubes of Smarties® or a similar coloured product.

Aim:

To illustrate the importance of number distributions and probabilities.

Procedure:

1. For each of the tubes, record the total number of sweets as well as the number of each individual colour. Smarties® are made as individual colours which are then mixed and the tubes filled by weight not by number. This means that certain colours will be missing and, because not all the Smarties® are the same weight, different tubes will often contain a different number of sweets.

2. If a new colour is introduced, what proportion must be used in the original mixture to ensure that there is at least one of the new type in each tube? What is the probability of a tube containing only one colour?

3. Plot the number distribution (histogram) for the contents, using the fact that there is a range of values for the total number of Smarties® within the tubes.

4. Plot a pie chart showing the percentage of each colour of Smarties® present (see Figure 12.6).

5. If, as a promotion, the marketing department offers 10% free, how many more Smarties® on average will you get? Is it possible to have 10% extra weight and yet have fewer Smarties®?

PROJECT 13: CHROMATOGRAPHY OF COLOURS

I thank Dr. Alberto Taboada for updating the experimental procedure.
Apparatus:

EYE AND HAND PROTECTION ARE ADVISED, AS CHEMICALS ARE INVOLVED.

Coloured sweets such as Smarties\textsuperscript{\textregistered} or M&M’s\textsuperscript{\textregistered}.

100 ml beaker.

Water bath.

Hot plate.

Watch glass

1\% ammonia solution.

1 m white wool.

Ethanoic (acetic) acid.

Ethyl acetate.

2-propanol (isopropanol).

Aim:

To separate a food colours into their individual components.

Procedure:

1. Take five different coloured sweets (say red, green, blue, brown and yellow).
2. Place the sweets in 15 ml hot water in a 100 ml beaker for five minutes and allow the surface and the colour to dissolve before discarding the centres.
3. Boil approximately 1 m of white wool for five minutes in 1% ammonia solution.
4. Rinse the wool in cold water and then immerse in the beaker with the dye extract.
5. Acidify the solution with dilute ethanoic acid and simmer for five minutes.
6. Remove the wool from the beaker and rinse in cold water.
7. Re-extract the dye from the wool by soaking in 10–15 ml 1% ammonia solution for five minutes.
8. Remove the wool from the ammonia solution.
9. Evaporate the ammonia solution to dryness on a water bath.
10. Add four drops of water to the extracted dyes and spot onto a TLC plate (Merck, aluminium sheet 2 m x 6 cm; 60 F254) and allow to dry thoroughly.
11. Pour a solvent made up of ethyl acetate (40 parts), 2-propanol (30 parts) and water (25 parts) into a beaker so that it is about 5 mm from the bottom. Put the TLC plate into the solvent with the spot about 10 mm up from the bottom of the beaker.
12. Place a watch glass on top of the beaker and wait until the solvent almost reaches the top of the beaker.
13. Remove the TLC plate. The colours should have separated into their components.

Changing the ratio of the solvent components will alter the degree of separation. This technique was initially developed by Nestlé Rowntree Customer Services Department for synthetic colours and may not work so well for some natural colours.

**PROJECT 14: EFFECTIVENESS OF DIFFERENT PACKAGING MATERIALS**

**Apparatus:**

Balance capable of reading to 0.1 g.
Desiccator.
Silica gel.
Aluminium or glass dishes about 5 cm in diameter and 1 cm deep. Wrapping material from confectionery, including aluminium foil, paper and plastics. Sealing wax or odour-free adhesive. White chocolate buttons. Peppermint oil.

**Aim:**
To assess the relative effectiveness of different packaging materials as barriers to moisture and odour transfer.

**Procedure:**

**Moisture Barrier**

1. Fill the base of several dishes with the same weight of silica gel.
2. Cut a circle of the packaging material, slightly bigger than the dish and seal it over the top using the wax or an adhesive. Be careful that there is a good seal all the way around. Then record the weight of each of the dishes.
3. Put water in the bottom of the desiccator and then place the dishes inside it as shown in Figure 12 7.
4. Take the dishes out of the desiccator every day and reweigh them.

The poor moisture barriers will pick up weight relatively quickly. Eventually the weight will become constant again, because the silica gel won’t be able to pick up any more moisture.

**Odour Barrier**

White chocolate is used as it has a relatively bland taste.
1. Place about six buttons in each dish, but this time have at least five dishes for each type of packaging material.
2. Seal the packaging material over the tops of the dishes. This time there is no need to weigh the dishes.
3. Put the peppermint oil in the base of the desiccator and then put the dishes on the tray.
4. At intervals (e.g. 1, 5 and 14 days), take out one dish for each type of packaging material. Get five different people to taste
the buttons and to score them for the intensity of the peppermint flavour on a scale of 1 to 10.
5. Plot intensity versus time for the different packaging materials.

**Leak test**

1. Place a flow-wrapped sample or bag under the surface of the water in a container and squeeze gently.
2. Count the rate at which bubbles come to the surface. If the rate is such that you can count aloud the number of bubbles, then the packaging is satisfactory even though some leaks still exist.

**PROJECT 15: VISCOSITY AND FLAVOUR**

**Apparatus:**

Bars of chocolate.
Refrigerator.
Knife.
2 pots of set yoghurt.

**Aim:**

To show that the speed at which a food melts in the mouth affects its taste as well as its texture. This is because the chocolate viscosity
affects the speeds at which the different molecules reach the flavour receptors (Figure 5.1).

**Procedure:**

1. Use the knife to scrape shavings of chocolate from one bar.
2. Place one bar and half the shavings in a refrigerator or freezer for 24 hours.
3. Keep the other half of the shavings with another bar of chocolate at room temperature.
4. Taste all four samples, recording their hardness, speed of melt, creaminess and cocoa intensity. Although all were originally the same, large differences should be recorded.
5. A similar effect can be obtained from set yoghurt. Stir one pot vigorously so that it becomes a thin liquid, then compare the taste of this with the other thick sample.

**PROJECT 16: HEAT-RESISTANCE TESTING**

**Apparatus:**

*Part a*

Oven set at 32°C.
Filter paper.
Several different brands of chocolate.
Knife.

*Part b*

Refrigerator.
Balance (accurate to three places of decimals).

**Aim:**

Chocolate melting and sticking to the packaging is a major problem in warm or hot climates. This experiment shows one method of measuring how badly this is likely to happen and that the chocolate type affects its heat resistance.
Procedure:

Part a

1. Mark the filter paper so that it is divided into small squares.
2. Cut the chocolate into equal sized squares.
3. Place the samples in the middle of the filter paper in the oven at 32 °C for two hours.
4. Remove each filter paper from oven and take the chocolate off it.
5. Count the squares that are fat stained.

The more squares that are affected the easier the chocolate melts. It may be possible to relate this to the smoothness of the chocolate. Why is this?

Part b

1. Break off four pieces from the bar.
2. Determine the weight of the four pieces.
3. Put two pieces one way up on the filter paper, the other two the other way up.
4. Place these in an oven at 32 °C for two hours.
5. Remove and put immediately in a refrigerator for at least one hour.
6. Remove the chocolate and reweigh it.
7. Determine the fat loss.
8. The effect of oven temperature, and chocolate size and shape can also be investigated.

PROJECT 17: COEFFICIENT OF EXPANSION

Apparatus:

Water bath.
Flask fitted with a stopper and with a glass tube through the middle (Figure 12.8).
Thermometer accurate to at least 0.5 °C.
Chocolate.
Sugar solution.
Aim:
Some chocolate confectionery products crack because two of the components expand at different rates when the temperature changes.

Procedure:
1. Melt the chocolate at about 40 °C for several hours, then cool to 38 °C.
2. Set the water bath to 38 °C.
3. Pour the chocolate into the flask so that it is almost full, and push the bung and tube into the top. (CARE: WEAR GLOVES IN CASE THE TUBE BREAKS.)
4. Place the flask in the water bath so that the neck is just above the water.
5. Allow the system to come to equilibrium and then measure the height of the chocolate up the tube from the bung.
6. Raise the temperature by 2 °C and leave for 20 minutes, then remeasure the height.
7. Repeat up to 50 °C.
8. Plot the height of the chocolate column against temperature.
9. Repeat the experiment but using sugar solution.

The two curves should be different.

PROJECT 18: THE MAILLARD REACTION

Apparatus:

Glucose.
Valine. (Can be purchased from national chemical suppliers.)
Small beaker (100 ml).
Hot plate.
Sunflower oil.

Aim:

To try to develop some of the chocolate aroma that is formed during roasting.

Procedure:

1. Dissolve about 3.6 g glucose and 0.6 g valine in 20 ml water.
2. Add about 2 ml sunflower oil.
3. Heat to near boiling point for about 15 minutes. (Try to simmer, and stir occasionally.)
4. Smell the aroma. CARE – VERY HOT – USE GLOVES AND SAFETY GLASSES, TAKE EXTRA CAUTION IF IT IS SPITTING.
5. Test for different combinations of temperature, time and relative concentration.

Valine is only one of the many amino acids that are present in cocoa, so it will not be possible to generate the full aroma of chocolate.
**Glossary**

**Acetic acid**: Common name for ethanoic acid.

**Chocolate crumb**: A dehydrated mixture of milk, sugar and cocoa liquor, used as an ingredient for some types of milk chocolate.

**Cocoa butter**: Fat pressed out from the centre (nib, cotyledon) of cocoa beans.

**Cocoa butter equivalent**: A fat that can be mixed with cocoa butter in any proportion without upsetting the way it crystallises.

**Cocoa liquor**: Cocoa nib which is finely ground. Like chocolate, it is solid at room temperature but liquid above 35°C.

**Cocoa mass**: Another name for cocoa liquor.

**Cocoa nib**: Cocoa beans with the shell removed.

**Conche**: A machine which mixes the chocolate ingredients to make it into a liquid and to remove some of the unwanted flavours.

**Enrober**: A machine which coats sweet centres by pouring chocolate over them.

**Ethanedioic acid**: Also known as oxalic acid.

**Ethanoic acid**: Also known as acetic acid.

**Lauric fat**: A fat rich in dodecanoic (lauric) acid (12:0). It is a major component of fats from coconut and palm kernel.

**Outer**: A box containing a number of bars or tablets.

**Oxalic acid**: Also known as ethanedioic acid.

**Phosphoglyceride**: A fat containing phosphoric acid (or other phosphorus-containing acids) in and appropriate ester form such as glycerophospholipid.

**Phospholipid**: Commonly used name for phosphoglyceride.

**Plastic viscosity**: A measurement related to the viscosity of a liquid when it is moving relatively quickly.
Polymorphism: The ability of a substance to crystallise in several different forms with different melting points.
Tempering: A process for ensuring that the fat in the chocolate sets in the correct crystalline form.
Triacylglycerol: A class of fats made up of glycerol esterified to three fatty acids.
Triglyceride: Commonly used name for triacylglycerol.
White chocolate: Chocolate made from cocoa butter, sugar and milk powder.
Yield value: A measurement relating to the energy required to start a liquid flowing, *i.e.* its viscosity when it is moving very slowly.
Subject Index

Acetobacter bacteria 20
acidity, conching 69
acne 203–4
Aero® 171, 190
After Eight® 139
air bubbles 134, 176–9
airflow and particle size 64
alcohols, sugar 28–9
alkalising process see Dutching process
allergens 185
allergies 204
amino acids 20
amorphous sugar 25–6, 209–10
Anna of Austria 2
anti-caries factors in cocoa 202
antioxidants 8, 9–10, 205–6
Aztecs 1
Bahia, Brazil 4, 13
ball fall viscometers 160–1, 214
ball mills for cocoa beans 53–4
beans see cocoa beans
behenic acid 117
Benefat™ 124
Bingham fluids, viscosity 83
biopolymers for packaging 193–4
bloom see fats bloom
body mass index (BMI) 200–1
Borneo 119
Bounty® 192
box fermentation 16, 16–17, 19
browning reaction 27–8
butter see cocoa butter
cacao tree see cocoa
Cadbury
flavour 37, 163
lecithin 97
marketing 9
Quakers 4
caffeine 206, 207
Caprenin 124
caramel 137–8
carbohydrates 198–9
Cardinal Richelieu 2
caseins 32–3, 202
Casson equation, viscosity 82–3, 88–9, 162
catechins 9–10
CBEs see cocoa butter equivalents
CBIs see cocoa butter improvers
cellulose based biopolymers 193–4
‘Charlie and the Chocolate Factory’ 125
CHD see coronary heart disease
chemical changes, conching 69–70
derelles 14–15
chocolate bloom 90
chocolate crumb 7–8, 36–7
cholesterol 198
chromatography
colours 223–5
flavour 164
cleaning, cocoa beans 39–40
cocao tree see cocoa
cocoa butter
coating
chocolate 147–50
conching 68–9
sugar cubes 86–7
Subject Index

235

cocoa
anti-caries factors 202
colour 3
countries producing 11, 12–13
fermentation 15–20
liqour 42–3
nibs 3, 8, 44–5, 49–51
pods 14–16, 17
powder 57
trees 11–12
see also cocoa butter
cocoa beans 14–15, 39–58
butter 54–7
chemical changes in roasting 46–7
cleaning 39–40
drying 20–2
Dutching process 3, 54–7
fermentation 16, 18–20
liqour 46, 50–1
Maillard reaction 47–9
mills 51–4
nibs grinding 49–54
nibs roasting 44–5
powder 57
roasters 45–6
roasting 40–3, 44, 46–7
size variation 40–3
storage 22–3
Strecker reaction 49
transport 22–3
winnowing 43–4
cocoa butter
crystalline forms 107–10
eating 3
fat 1
fat phase 173–4
lauric fat 121–2
melting profiles 114
non-lauric fat 122–4
polymorphic forms 109–10
pre-crystallisation 110–12
production 55–7
replacers 121–2
separation 213–14
solid fats 113, 123
structure 104–7
temperature ranges 107–8
cocoa butter equivalents (CBEs) 118–19
cocoa butter improvers (CBIs) 120
Cocoa Tree Chocolate House 2
cocnut oil 121
Codex Alimentarius 185
coefficient of expansion 148–9, 229–31
colours
chromatography 223–5
cocoa 3
Columbus, Christopher 1
combined milling 60, 64–8
composition of chocolate 222
conche machines 7
conching 68–79
acidity 69
chemical changes 69–70
Frisse machines 77
machines 7, 59, 60, 73–8
moisture 69
physical changes 70–1
power 100–1
stages 78–9
viscosity 71–2
conduction 144
confectionery, magnetic resonance imaging 168
continuous low volume conching machines 76–8
continuous phase experiments 218
convection 144
coolers 145–6
cooling curves for chocolate 130–1
coronary heart disease (CHD) 197–8
cotyledons see nibles
countries producing cocoa 11, 12–13
cream 92
cream eggs 180–3
Criollo cocoa 11, 13
Crunchie® 125, 139, 222
crystal packing of triglycerides 108
crystalline sugar 24, 26, 209–10
crystallisation
cocoa butter 107–10
cocoa butter temperatures 108
measurement 167–70
triglycerides temperatures 106
crystals, differential scanning calorimetry 169–70
Subject Index

Dairy Milk® 192
dairy processes 30, 34
Delta chocolate 6
Desert Bar™ 176
diabetic chocolate 172
differential scanning calorimetry 169–70
disc mills for cocoa beans 52–3
distribution of Smarties® 223
Don Cortez 2
double chain packing of fats 107
Drifter Bar® 222
dry conching 78
drying of cocoa beans 20–2
Dutching process 3, 54–7

E numbers 185
Easter eggs 136, 190
Ecuador 11
elaidic acid 122
emulsifiers, viscosity 93–9
enrobers 59, 125, 139–42
enzyme interesterification 120
enzymes in fermentation 18
epicatechin 55
equilibrium relative humidity (ERH)
  shelf life 186–7
  sugar 26–7
  tempering 126
European Union (EU) legislation 184–5
eutectics, fats 112–16

fats
  additions and viscosity 90–1
  bloom 31, 104, 109–10, 116–18, 144, 173
  chain packing 107
  cocoa powder 57
  eutectics 112–16
  measurement 158–9
  migration 117–18, 212–13
  migration and shelf life 186
  mixing 112–16
  nutrition 197–8
  phase 173–4
fatty acids 69
  ‘feet’, liquid chocolate 60
fermentation
  box 16–17
  cocoa 15–20
  enzymes 18
  heap 15–16
  pH 19
  Fernando Po 4
  fine ingredient milling 60
  fineness of particles, viscosity 88–9
  flavanols 204–6
  flavonoids 9–10
  flavour
    analysis 163–5
    conching 68, 69
    house 37
    receptors 80–1
    viscosity 227–8
flow cap viscometers 160–1
flow of chocolate, moisture 92–3
flow wrap 190–3
foil wrap 189–90
Forastero cocoa 11, 13
force v. distance curves in three point
  bend tests 166–7
Fox’s Glacier Mints® 209
Fraunhofer diffraction 155
Frisse conche machines 75–7
fructose 23, 28
Fry, Joseph 4
galactose 26
ganache 92
gas, air bubbles 178
Ghana (Gold Coast) 4, 13, 21
GI see glycaemic index
gloss, chocolate coating 149–50
glossary 232–3
glucose 23, 26, 28
glycaemic index (GI) 199
grinding of cocoa nibs 49–54
HACCP see hazard analysis of critical
  control points
hand tempering 128–9
hand-decorating tools 141–2
hard-coating process 151–2
hardness 165, 167, 220–1
hazard analysis of critical control points
  (HACCP) 40
hazelnuts 117
HDLs see high density lipoproteins
Subject Index

headaches 203
heap fermentation 15–16
heat-resistance testing 228–9
Hershey
  Desert Bar™ 176
  flavour 37, 163
Quakers 4
high density lipoproteins (HDLs) 198
high pressure liquid chromatography (HPLC) 41, 164–5
high-cocoa chocolate 172
hollow eggs 137
HPLC see high pressure liquid chromatography
ice-cream coatings 172–3
illipe nuts 119
impact mills for cocoa beans 52
indices for solid fats 115
Indonesia 13
invert sugar 23
Ivory Coast 13
‘joccalatte’ chocolate drink 2
Karl Fischer reaction 92, 157
KitKat® 9, 174, 190, 191

lactitol 28
lactose 26–8, 36
lauric fat, cocoa butter 121–2
LDLs see low density lipoproteins
leaking chocolate centres 182
lecithin
  effects 217–18
  labels 185
  mixing 100
  viscosity 94–8, 100
legislation 184–5
Lindt, Rodolpho 6
Lindt, Rudi 68
Lion Bar® 125, 186, 222
liquid chocolate 59–61
  conching 78–9
  ‘joccalatte’ 2
  mixing 99–101
  mouth 80
  particles size 80–1, 84–90

storage 125–6
  viscosity 81–4, 86–90, 100–1
liquor
  cocoa beans 50–1
  roasting 44–5
long conche machines 73–4
Louis XIII 2
low calorie chocolate 171
low calorie fats 124
low density lipoproteins (LDLs) 197, 205
M&M’s® 146, 151
machines, tempering 126–8
magnetic resonance imaging (MRI) 168–9
Maillard reaction 27–8, 37, 47–9, 70, 231
Malaysia 13, 119
mannitol 28
marketing in UK 9
Mars Bar® 9, 125, 139, 174, 222
Maya people 10
melting profiles of cocoa butter 114
meters for tempering 130–2
methylxanthines 206–7
migraine 203
migration of fats 117–18
milk 30–6
  fat 31–2, 114–15
  fatty acids 32
  powders 33–4, 36
  proteins 32–3
  tooth–friendly proteins 202–3
milk chocolate 59–60
  milk fat 114–15
  particle size distribution 155–6
  solid particles 88–9
  viscosity 134–5
Milkybar 190
milling
  chocolate 61–8
  cocoa nibs 51–4
  combined 64–8
  separate ingredient grinding 62–4
mixing
  fats 112–16
  liquid chocolate 99–101
moisture
  analysis 156–8
  conching 69
flow of chocolate 92–3
migration and shelf life 186
Montezuema 1
moulding
shells 135–9
single-shot depositors 181
solid tablets 132–5
mouth, liquid chocolate 80
MRI see magnetic resonance imaging
mucor miehei enzyme 120–1
multiple chocolates centres 183

Nacional cocoa 11
Nestle, Henri 4
Newton, Isaac 82
Newtonian liquids, viscosity 82–3
nibs
  grinding of cocoa beans 49–54
  milling 51–4
  roasting of cocoa beans 44–5
Nigeria 13
NMR see nuclear magnetic resonance
no added sugar chocolate 171
non-cocoa vegetable fats 118–24
non-enzymic browning see Maillard reaction
non-lauric fat replacers for cocoa butter 122–4
nuclear magnetic resonance (NMR) 167–9
nutrition 196–9
nuts in chocolate centres 182

obesity 200–1
oleic acid 105, 119, 122–3
oral clearance, tooth decay 203
oxalic acid, tooth decay 203

packaging 187–95
  biopolymers 193–4
  effectiveness 225–7
  flow wrap 190–3
  foil wrap 189–90
  paper wrap 189–90
palm kernel oil 121
palmitic acid 105, 119
panning
  coating centres 125, 146–52

rotational movement 147–8
sugar 150–2
surface curvature 147–8
paper wrap 189–90
particles
  diameter 85
  fineness and viscosity 88–9
  separation 210–12
  size 62–4, 80–1, 84–90, 216–17
  size measurement 153–6
  specific surface area 88
  viscosity 86–90
see also solid particles
paste phase conching 78
peanuts 185
Pepys, Samuel 2
peroxides in milk fat 31–2
Peter, Daniel 4–6
petrol stations 163
PGPR see polyglycerol polyricinoleate
pH
  Dutching 55
  fermentation 19
  teeth 202
phenylethylamine 207
physical changes in conching 70–1
PLA see polylactic acid
Plantic, biopolymers 194
platelets 205
Polos® 209
polydextrose 29–30
polyglycerol polyricinoleate (PGPR) 98–9
polyhydroxyl polyesters 193–4
polylactic acid (PLA) 193–4
polymorphic forms of cocoa butter 109–10
polyols see sugar alcohols
polydextrose 29–30
powders
  cocoa 57
  lactose 36
  milk 33–4
  whey 36
  pralines 117
pre-crystallisation of cocoa butter 110–12
pressure in air bubbles 177
procyanadins 204–6
proteins
  milk 32–3
  nutrition 199
psychoactive compounds 206–8
pulsed nuclear magnetic resonance 168
pyrazines 164–5
Quakers 4
radiation 144
raisins in chocolate centres 182
rationing of chocolate in UK 9
reduced/low fat chocolate 172
relative humidity see equilibrium
  relative humidity
replacers for cocoa butter 121–2, 122–4
roasting
  bean size 40–2
  beans 44
  chemical changes 46–7
  Maillard reaction 47–9
  nibs 42–3
  roasters 45–6
  temperature 41–2
  whole bean 42–3
robotic packaging 194–5
roll refiners 64–7
Rolo® 190
rotary conches 74–5
rotating drums 146–7
rotational movement in panning 147–8
Rowntree 4, 118
sal crop 119
Salatrim 124
Sanders, Nicholas 2
sealing in flow wrap 192–3
separate ingredient grinding mills 62–4
shape, chocolate coating 148
shape-retaining chocolate 173–6
shea crop 119
shear
  pre-crystallisation rates 111–12
  viscosity 71–2, 82
  viscosity measurement 162–3
shelf life 186–7
shellac 150
shells 135–9
single-shot depositors 180–1
size
  air bubbles 178–9
  cocoa beans 40–3
  distribution in liquid chocolate
    particles 84–5
Smarties® 9, 146, 151, 188–9, 223
‘smoky’ flavour of chocolate 164–5
snap, chocolate 165
solid fats
  cocoa butter 105, 113
  cocoa butter replacement 121–3
  indices 115
solid particles
  framework of chocolate 175–6
  milk chocolate 88–9
  vibration 134–5
  solid tablets in moulding 132–5
  solidification of chocolate 143–6
  Sollich Solltemper tempering machine 126–7
SOO triglycerides 105–6
sorbitol 28
SOS triglycerides 105–6
soxhlet method for fat content 158–9
soya lecithin 97
specific surface area of particles 88
spheres packing 87
spinning in hollow eggs 137
SSS triglycerides 104–6
standard method for viscosity 161–2
starched based biopolymers 193–4
stearic acid 105, 119
sticking of sweets 182
storage of cocoa beans 22–3
Strecker reaction 49
sucrose (saccharose) 23–4
sugar
  alcohols 28–9
  amorphous 25–6, 209–10
  beet 23–4
  cane 23–4
  chocolate 1, 23–30
  cooling effect 29
  crystalline 24, 26, 209–10
cubes coating 86–7
panning 150–2
Subject Index

polydextrose 29–30
sugar free chocolate 171
sweetness 29
surface active agents 95–6, 98–9
surface curvature in panning 147–8
‘sweet chocolate’, USA 185
sweetness of sugars 29
sweets, sticking 182

TA texture measuring instrument 165–6
‘tails’ 137–8, 142–3, 182
tannins 55
temperature
  crystalline cocoa butter 107–8
  roasting 41–2
  roll refiners 67–8
  shapes 181
  solidification 144
  tempering 131–2
  viscosity 128
tempering
  chocolate 142–3
  hand 128–9
  liquid chocolate storage 125–6
  machines 126–8
  measurement 128–32, 219
  meters 130–2
  temperature 131–2
  see also pre-crystallisation
texture monitoring 165–7
theobromine 206
three-point bend tests 165–7
Toblerone® 188–9
tooth decay 201–3
tooth-friendly milk proteins 202–3
transparent coatings 174
transport of cocoa beans 22–3
trees, cocoa 11–12
triglycerides
  cocoa butter 104–5
  crystal packing 108
  crystallisation temperatures 106
  milk fats 31
Trinitario cocoa 11
triple chain packing of fats 107
Turkish Delight 188
Twix® 192
Unilever 118, 120
United Kingdom (UK) marketing 9
United Nations 185
USA legislation 185
vacuum formed trays (VFTs) 192
Van Houten 3
VFTs see vacuum formed trays
vibration 134–5
viscosity
  Bingham fluids 83
  caramel 137–8
  Casson equation 81–2, 88–9, 162
  chocolate centres 182
  cocoa beans liquor 50–1
  conching 71–3
  emulsifiers 93–9
  fat addition 90–1, 214–16
  fineness of particles 88–90
  flavour 227–8
  lecithin 95–6
  liquid chocolate 59, 81–4, 86–90, 100–1
  measurement 159–62
  milk chocolate 134–5
  moisture addition 214–16
  Newtonian liquids 82–3
  shear stress 162–3
  temperature 128
  units 82–3
  viscometers 160, 161–2

water
  evaporation bubbles 179
  shape 174–5
  ‘waterfalls’ 125, 139–41
  weight control in chocolate production 222
whey 32–3, 36
white chocolate 8–9
White’s Chocolate House 2
whole bean roasting 42–3
winnowing, cocoa beans 43–4
witches’ broom disease 13
xylitol 28

Yucatan 1–2
Ziegleder, Dr. 70