Animal skin that has been processed to retain its flexibility, toughness, and waterproof nature is known as leather, and the leather industry is New Zealand's seventh largest industry. New Zealand leather is made by processing the skins of animals slaughtered in the meat processing industry, and hence the majority of skins tanned are those of sheep and cattle. Animal skins are converted to leather in an eight step process as follows:

**Step 1 - Unharing**
The animal skins are steeped in an alkali solution that breaks down the structure of the hair at its weakest point (the root) and so removes the hair.

**Step 2 - Liming**
The hairless skin is immersed in a solution of alkali and sulphide to complete the removal of the hair and to alter the properties of the skin protein (collagen). The collagen becomes chemically modified and swells, leaving a more open structure.

**Step 3 - Deliming and Bateing**
The skin structure is then opened further by treatment with enzymes, and further unwanted material is removed.

**Step 4 - Pickling**
The skins are then treated with acid to preserve them for up to two years.

**Step 5 - Tanning**
This is the most chemically complex step. During tanning, the skin structure is stabilised in its open form by replacing some of the collagen with complex ions of chromium. Depending on the compounds used, the colour and texture of the leather changes. When leather has been tanned it is able to 'breathe' and to withstand 100°C boiling water, as well as being much more flexible than an untreated dead skin.

**Step 6 - Neutralising, Dyeing and Fat Liquoring**
The leather is then treated with alkali to neutralise it and so prevent deterioration, and then dyed. This involves fixing a variety of compounds onto the chromium, as that is the most reactive site present. Once the leather is dyed, it is treated with reactive oils that attach themselves to the fibrous structure, improving suppleness and flexibility.

**Step 7 - Drying**
Water is removed from the leather, and its chemical properties stabilised.

**Step 8 - Finishing**
A surface coating is applied to ensure an even colour and texture, and to improve its ability to wear. Suede leather is also buffed at this point to give it its distinctive finish.

Animal skins that are processed in New Zealand go on to be made into a variety of leather goods, or are exported in an unfinished condition to be further treated overseas.
INTRODUCTION

When an animal is alive, its skin is soft, flexible, very tough and hard wearing: it has the ability to allow water vapour to pass out, but it will not allow water in. When the skin dies it loses these characteristics: if it is kept wet it rots, and if it is dried it goes hard and brittle. The process of tanning is to retain the skin's natural properties, to stabilise its structure and at the same time to chemically process it so it will no longer be subject to putrefecation. Thus leather is animal skin that has been treated such that its natural properties are retained.

Skin is made up of many bundles of interwoven protein fibres which are able to move in relation to one another when the skin is alive. When the skin dies, these fibres tend to shrivel and stick together. Essentially, the purpose of tanning is to permanently fix the fibres apart by chemical treatment, and to lubricate them so they can move in relation to one another. Well tanned leather, therefore, retains the properties of flexibility, toughness and wear. It also continues to 'breathe', allowing water vapour to pass through but remaining reasonably water-proof. It is this characteristic which accounts for the comfort of genuine leather shoes and clothing.

In addition, the process of tanning imparts the advantage of resistance to heat. This is an important factor in many of the uses of leather. In conjunction with chemical processing, the tanner imparts colour, texture and finish to the leather, to enhance its appearance and suit it to today's fashion requirements.

The basic component of the skin is collagen, a fibrous protein. The latest research indicates that the basic collagen structure consists of twined triple units of peptide chains of differing lengths. The amino acid residues are joined together by peptide links. The peptide chains within the triple helices are held together by hydrogen bonding (Figure 1).

The main amino acid components of skin are given in Table 1.
Table 1 - Amino acids found in skin

<table>
<thead>
<tr>
<th>Amino acid</th>
<th>Structure</th>
<th>Abundance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glycine</td>
<td><img src="image" alt="Glycine Structure" /></td>
<td>26.8% of total nitrogen</td>
</tr>
<tr>
<td>Alanine</td>
<td><img src="image" alt="Alanine Structure" /></td>
<td>8.0%</td>
</tr>
<tr>
<td>Proline</td>
<td><img src="image" alt="Proline Structure" /></td>
<td>9.0%</td>
</tr>
<tr>
<td>Hydroxy proline</td>
<td><img src="image" alt="Hydroxy Proline Structure" /></td>
<td>8.0%</td>
</tr>
<tr>
<td>Arganine</td>
<td><img src="image" alt="Arganine Structure" /></td>
<td>15.3%</td>
</tr>
</tbody>
</table>

There has been no conclusive evidence showing that the amino acid residues are arranged in any particular order within the peptide chain.

Skin collagen is usually associated with keratin (the protein in hair, wool and nails). Most mammals have an outer coat of hair, wool or fur, which acts as an insulating layer and keeps the animal warm. Keratin is a fibrous protein and different from collagen in one very important aspect: the polypeptide chains are linked together by cystine linkages.

The sulphur-sulphur linkage in cysteine is susceptible to the action of alkali, and breaks down quite readily in the presence of alkali and a reducing agent.

The early part of leather production is the removal of hair from the skin, and the presence of the cystine linkage makes it possible for this to be achieved.
Physically, the skin consists of tiny interwoven collagen fibres, each of which in its turn is made up of tiny fibrils. The tanner aims to separate these fibres, remove any unwanted matter from the structure and stabilise and preserve it, whilst retaining the useful properties.

**Raw materials**
The basic raw material for the New Zealand tanning industry is a by-product of the meat processing industry. Approximately 25 million lambskins, 8 million sheepskins, 2.2 million cattle hides, 1.2 million calfskins, 200,000 goat skins and 400,000 deerskins are available each year for processing into finished leather products. Cattle hides give leather for shoes (soles and uppers), clothing, belts and upholstery. Calfskins are used for fashionable shoe uppers, gloves and for clothing leathers. Sheepskins and lambskins with wool on are used for car seats, floor rugs, clothing, bedding and footwear, while, with the wool removed, they are used to produce suede and grain clothing leathers as well as chamois leathers.

The tanning materials used in the conversion of skin to leather are mostly imported. There are four main types:

- Vegetable - extracts from the bark and wood of trees (oak, wattle etc.).
- Mineral - in the main, trivalent chromium sulphate. This is produced by the reduction of hexavalent chromium in the form of sodium chromate or dichromate by organic or inorganic reducing agents.
- Aldehydes - formaldehyde, glutaraldehyde or oxazolidine
Synthetic replacements for the vegetable tanning materials.

THE TREATMENT PROCESS - PREPARATION FOR TANNING (Steps 1 - 4)

The main processing sequences of leather manufacture are shown in Figure 2. Where skins are processed with the hair or wool on, the unhairing and liming processes are omitted and replaced by a scouring (washing) of the wool or hair. The main chemical processes carried out by the tanner are the unhairing, liming, tanning, neutralising and dyeing.

Step 1 - Unhairing
The keratin of hair and wool is subject to attack by alkali, which will break (by hydrolysis) the sulphur-sulphur bond in the cystine linkage of the keratin. The hair is attacked first at the root, where it is in its immature form. After a certain time of being steeped in a solution of alkali (sodium or calcium hydroxide) and a reducing agent, normally sodium sulphide, the hair roots are dissolved, and the hair may be removed and saved.

In the processing of sheep and lambskins, the solution of alkali and sulphide is applied on the inside of the skin, allowing the solution to penetrate through the skins, attacking the wool roots, which can then be removed by 'pulling' them from the skin. This is the origin of 'slipe wool', which forms a sizeable portion of New Zealand's wool production.

Step 2 - Liming
The process of liming is a combination of chemical and physical action on the skin structure. The chemicals involved are similar to those in the unhairing operation. A solution of lime and sodium sulphide is used and the skins are immersed in this solution for whatever time is necessary to produce the desired effects. The process of unhairing is taken to completion during the liming process, and there is appreciable modification of the collagen due to the action of alkali.

\[
\text{Skin protein (collagen)} \rightarrow \text{alkali} \quad 1) \text{Hydrolysis of amide groups} \\
2) \text{Modification of guanide groups} \quad 3) \text{Hydrolysis of keto-imide links in protein chains} \\
4) \text{Swelling} \quad 5) \text{Removal of unwanted material}
\]

\[
\text{Hydrolysis of amide groups} \\
R\text{C}==\text{NH}_2 + \text{OH}^- \quad \rightarrow \quad R\text{C}==\text{O}^- + \text{NH}_3
\]
Waste Products | By-products | Process Chain
--- | --- | ---

| Hides & skins | | |
| Salts | | Soaking | Prepared skins |
| | | | |
| Excess alkali and sulphide | Hair or wool | Hair removal | Liming | Unhaired skins |
| Acid | | | Pickling | |
| Chromium salts | | | Chrome tanning | Skins ready for tanning |
| Acid and dyes | | | Neutralizing and Dyeing | Dyed leather |
| Dust | | | | Drying | Dried leather ready for finishing |
| | | | | | Surface coating |
| | | | | Buffing | Finished leather |
| | | | | | |
| | | | | | Finished suede Clothing manufacturers Bag manufacturers |
| | | | | | Car and Furniture manufacturers |
| | | | | | Clothing manufacturers Shoe manufacturers Shoe Industrial uses manufacturers |

Figure 2 - Schematic of the treatment process
2) Modification of guanide groups (arginine residue)

\[
R-(CH_2)_3-NH-C-NH_2 + H_2O + OH^- \rightarrow R-(CH_2)_3-NH-C-O^- + 2NH_3
\]

3) Hydrolysis of amide linkages in the backbone of the collagen chains

\[
\text{CH}_2-C-NH-C\text{-CH}_2 + OH^- \rightarrow \text{CH}_2-C-O^- + H_2N-C\text{-CH}_2
\]

4) Swelling

In addition to the chemical action, in the presence of an alkali, swelling of the fibrous structure takes place. This is due to an osmotic pressure effect. The result is the separation of the fibres and the fibrils from one another and an opening up of the whole structure.

5) Removal of unwanted material

Present in the collagen structure are globular proteins and other interfibrillary substances. These are hydrolysed and removed in the subsequent washing and bateing processes. Unwanted fats are saponified.

**Step 3 - Deliming and Bateing**

After the strong alkaline action, the skin structure is further opened up during the deliming and bateing process.

Bateing is an enzyme digestion process which continues the cleaning up process begun by the alkaline action. Bateing is carried out at pH 9-10. This is achieved by treatment with acid salts (ammonium chloride or ammonium sulphate) or carbon dioxide until the desired pH is reached.

Hydrolysis is continued by the enzyme process and further dissolution of unwanted material occurs. The main fibrous network is then cleaned up and the swelling is reduced. (See article on Fellmongery for more details)

**Step 4 - Pickling**

The main acidification process is that of pickling. The skins are agitated in a solution of salt and sulphuric acid until they are at or near equilibrium at a pH value of 3.0 - 3.5. (See article on Fellmongery for more details)

**THE TREATMENT PROCESS - TANNING (Step 5)**

The tanning or stabilising process can be carried out as noted before, by means of vegetable, mineral or synthetic tanning materials. The vegetable and synthetic tanning materials are largely anionic in nature and are essentially similar in their action on skin protein. Chemically they are complex mixtures and their chemistry is not well understood. The vegetable tans fall into two main classes.

- Hydrolysable tans - complex esters of phenol and carboxylic acids and polyhydroxyphenols
• Condensed tannins - mixtures of products of varying degrees of polymerisation, containing several molecules of catechin or catechin-like compounds

Syntans are largely condensation products of sulphonic acids (phenolic and naphtholic) and formaldehyde and similar general types.

Vegetable and synthetic tanning materials are widely used to modify the properties of chrome tanned leather, and as self tannages to produce sole leather, belting, football covers and other types of leather where stretch and suppleness in the leather is not required.

The main mineral tanning material is trivalent chromium sulphate. This is now widely used in the preparation of shoe upper leathers and clothing leathers, in fact, in all leathers where softness, stretch and suppleness are required. It is synthesised as follows:

$$\text{Na}_2\text{Cr}_2\text{O}_7 + 3\text{SO}_2 + \text{H}_2\text{SO}_4 \rightarrow \text{Na}_2\text{SO}_4 + \text{H}_2\text{O} + \text{Cr}_2(\text{SO}_4)_3$$

**The use of chromium in tanning**

Chromium, a transition element which forms co-ordination complexes by using 3d orbitals to accommodate extra electrons, has an unrivaled position as a tanning agent. The complex is basic by nature and will contain hydroxyl groups associated with the chromium nuclei.

There are two main features of chromium chemistry which enable it to act as a tanning agent. Firstly, the complexes formed are of intermediate stability, and thus exchange of co-ordinating ligands can take place comparatively easily. Secondly, chromium has the ability to form polynuclear complexes in which Cr-O-Cr bridges are involved. These come about as follows:

$$2\text{Cr(H}_2\text{O)}_6^{3+} \rightarrow 2\left[\text{Cr(H}_2\text{O)}_5\text{OH}\right]^{2+} + 2\text{H}^+ \leftrightarrow [\text{H}_2\text{O)}_4\text{Cr-O-Cr(H}_2\text{O)}_4]^4+ \rightarrow [\text{H}_2\text{O)}_4\text{Cr-O-Cr(H}_2\text{O)}_4]^{2+}$$

**Basic salt**

**Oxo compound**

**Oxo compound**

The final process, that of oxolation, is a slow one and is not readily reversed. In a similar manner, long chains may be built up. By means of these polynuclear units the gaps between collagen chains in the skin structure can be bridged, giving the tanning action.

Individually these two properties are found in the chemistries of many other elements, and do not give any great tanning ability. But the two properties together give great tanning potency.

Chromium shows a strong tendency to form coordination compounds with molecules containing carboxyl groups. The order of stability of complex ions involving chromium with other differing ligands is:
NO₃⁻ < Cl⁻ < SO₄²⁻ < H₂O < SO₃²⁻ < HCO₃⁻ < CH₂CO₂⁻ < (1) <
CO₂—(CH₂)₃—CO₂⁻ < (2) < CO₂—CO₂⁻ < CN⁻ < OH⁻

The chromium-collagen complex lies approximately between (1) and (2) in the range of stabilities, and the order in the series has a number of practical and theoretical implications. The process of chrome tanning is now thought to involve the ligand present in the chromium complex being replaced by collagen.

The chromium-collagen compound
There are a number of ways in which the tanning action could take place, and although an ionic compound may be formed initially, it seems certain that the main chemical action is by means of coordination complexes or covalent bonds. It also seems certain that the most important coordinators are the free carboxyl groups of the collagen structure, but the amino groups and the imido groups of the peptide linkage are not excluded. This theory is implied mainly by tanning actions at various pH's. Tanning action increases as pH increases, and no tanning occurs at all when all —CO₂— groups of the collagen molecules are protonated.

These factors are discussed below.

The nature of the anion
Increasing ability to tan: Cr(NO₃)₃ < CrCl₃ < Cr₂(SO₄)₃

The anion associated with the chromium salt is involved in the bridge between chromium atoms. SO₄²⁻ has the greatest ability to form this bridge.

The basicity of the chromium salt
At approximately 30-40% basicity, i.e. in the pH range 3.0 - 3.4, the average molecular size is approximately 2 - 4 chromium nuclei. This appears to be the optimum for tanning. At lower basicities the molecular size is too small and the complexes cannot bridge the gap between the collagen fibres, and at higher basicities the molecular size is such that the semi-colloidal particles cannot enter into the fibrous collagen structure.

pH value
The pH of a solution is determined by considering the difference in acidity of a solution at a given time and the equilibrium condition. Increasing pH gives greater tanning action.
Neutral salts
These also have an effect on the tanning action but this is not of great significance.

Complexing and masking agents
These are of major significance, and the type of leather produced depends very strongly on which complexing agents are present. These tend to slow down the tanning action. In some cases, e.g. that of oxalate, the ligand cannot be replaced by collagen, and hence a solution containing more than two molecules of oxalate to one of chromium will not tan. A solution of sodium oxalate will remove the chrome from chrome tanned leather. From the practical point of view the important complexing agents are sulphate, formate, acetate, phthalate, sulphite and the dicarboxylic acids. Sulphate is of low stability (hence the use of chrome sulphate powders). Formate is monodentate, is widely used, and slows the tannage down just enough to give the finished leather certain desired qualities. Acetate has an important colour effect, producing blue leather. Phthalate is bidentate (it occupies two coordination sites in the chromium nucleus). The use of this ligand increases the uptake of chromium, as it leaves fewer sites for coordination with the collagen substrate.

Practical aspects
How does this affect us in practice? It can be seen that in order to penetrate into the fibrous structure of the hide or skin, the chromium complex must be small. This may be obtained by a low pH and a low basicity chromium salt. When penetration has been achieved a higher pH is required; this liberates the carboxyl groups of the collagen and leaves them free for coordination, and helps in the formation of \( \text{\textit{oLo}} \) and \( \text{\textit{oxo}} \) polynuclear chromium complexes. When some tanning has been achieved and the skin is more resistant to heat, the temperature may be raised.

Practical tanning therefore involves bringing the skin to be tanned to an acid condition, obtaining penetration of chrome at low pH, raising the pH, and finally heating to obtain maximum fixing of chrome.

Two examples of how this goes on are as follows:

Acetate masking

The leather produced here is flat and supple, not particularly high in chrome content and pale blue green in colour (suitable for bleaching to white leather). Where acetate is used, single links between chains occur, the leather is kept flat and the chrome content is not very high.
Phthlate masking

This leather is fuller, not so flat, has a higher chrome content and is green in colour. Here, because of the bidentate nature of the phthalate as a complexing agent, the chromium uptake tends to be greater and there are more links within the individual chains.

The practical measure of degree of chrome tannage is that of thermal stability. At the completion of any tannage the leather is tested to see what temperature, under wet conditions, it will stand. If it will resist 100°C (boiling water) it is considered to be satisfactorily tanned.

**THE TREATMENT PROCESS - AFTER TANNING (Steps 6 - 8)**

**Step 6 - Neutralising, Dyeing and Fat Liquoring**

After tannage, leather must be neutralised to remove unwanted acids to prevent deterioration during the drying process, and to prepare the leather for the next stages in processing: dyeing and fat liquoring. Neutralising is often carried out using mild alkalis which have some effect on the chromium complex (sodium formate is a good example).

Next is dyeing, using a variety of dyestuff types. Commercial dyes are essentially mixtures of chemical compounds which can be fixed (combined) to the material to be dyed and which have appreciable resonance within the molecule giving colour. A simple example is Orange II:
Its colour comes from the presence of the hydroxide group in conjunction with the azo group, allowing a high degree of resonance. It can be fixed to the material to be dyed by means of the sulphonic acid group.

In chrome tanned leathers the most reactive sites are the chromium nuclei, and these form the main points of fixation for dyestuffs. As with other processes, the main factors controlling dyeing are the physical one of locating the dye in the required position within the leather structure and the chemical one of controlling the conditions so that reasonable penetration and fixation are achieved.

A wide range of dyes are used in leather processing and some examples are:

**Direct dyes:**

\[
\text{NaSO}_3 - \text{N}=\text{N} - \text{NH} - \text{CO} - \text{NH}_2 \]

*C. I. Direct red 118*

**Reactive dyes:**

\[
\text{Dye} - \text{NH} - \text{C} = \text{N} - \text{C} = \text{N} - \text{Cl} \]

*I.C.I. Procions*

**Pre-metallized dyes:**

\[
\text{SO}_3^- - \text{N}=\text{N} - \text{O} - \text{Cr} - \text{O} - \text{SO}_3^- \]

*C. I. Acid Blue 158*
Acid dyes:

\[ \text{Polar Brilliant Red 3B} \]

The growing awareness of the effects on health of prolonged exposure to certain industrial chemicals has led to the prohibition of the use of certain dye stuffs - examples are direct dye-stuffs based on benzidine. Consumer requirements for both wash and light fastness in leather goods has lead to a shift away from the more level dyeing direct and acid dyes to the faster pre-metallized and reactive dyes. Dyeing technology has had to keep up to improve the level dyeing properties of these dyes.

Fat liquoring is the process in which 'tanned' fibres are treated with reactive oils, which attach themselves to the fibrous structure, and lubricate them so that they can move readily in relation to one another, producing a soft, supple leather.

Step 7 - Drying
Finally, as water is removed from the system, the chemical condition is stabilised and the final properties of the leather are determined.

Step 8 - Finishing
Finishing consists of applying a surface coating - pigments or dyes bound in an organic (acrylic, butadiene or polyurethane) or protein (casein) medium. This enhances the natural qualities of the skin and covers such defects as scars, horn damage, seed scars etc., as may be present. In the case of suede leather, the suede is buffed to prepare a 'nap' and to produce a generally even texture. The main requirements for finishing are evenness and the reproducibility of colour and adequate wear and feel properties.

THE INDUSTRY

The leather industry in New Zealand is an integral part of our agricultural economy.

New Zealand's largest industry, the meat industry, has the hides and skins of the slaughtered animals as one of its by-products. The leather industry, New Zealand's seventh largest, has exports of $750 million (1994). About half this value is in raw hides and skins and the remainder in fully or partly processed leather or leather goods.

Levels of processing
Most sheep and lambskins are fellmongered (dewooleed, limed, bated and pickled in the meatworks or independent fellmongeries). Some of these (approx 8 million per year) are further processed into chrome tanned leather (wet blue) or into finished garment leather.
within New Zealand. The remainder - about 25 million - are exported in pickled condition.

About 2.5 million lambskins per year (some from imported raw skins) are tanned for wool-on lambskins used for floor rugs, babycare rugs, car seats, bedding, clothing and footwear. These are largely exported.

Of New Zealand's 2.2 million cattle hides per year, about 200 000 are exported in raw salted condition, 900 000 are chrome tanned and exported in the wet blue condition and the remainder are fully processed into finished leather which is used in New Zealand and overseas for shoe making and upholstery of cars and furniture.

Approximately 1.2 million calfskins are produced each year. These are exported in wet blue or raw salted condition.

Most of the 400 000 deerskins produced each year are converted into suede or nappa clothing leather and are exported as finished leather or as finished leather garments.

**Economic aspects**
The total value of exports from the leather industry for the year to 30 June 1994 was $750 million of which -

- $450 million was in raw hides and skins
- $80 million was in finished woolskins
- $58 million was in finished bovine leathers
- $40 million was in finished lambskins
- $98 million was in wet blue hides
- $29 million was in wet blue lamb pelts

The industry is supported by its trade organisation, the New Zealand Tanners Association, and its research body, the New Zealand Leather and Shoe Research Association and the Tanning Industry Training Organisation.

There are 16 tanning plants in New Zealand (in addition to freezing works and fellmongeries) and a total of 1800 people are employed.

The industry has a set target of $1 billion in exports by the year 2000. To do this, it will need to carry out considerably more added value processing than it does in 1995.

The exports from New Zealand tanneries are sent to all parts of the world with Asian markets showing the fastest rate of growth.

A significant contribution is made to New Zealand's economy by the tanning industry.

**ENVIRONMENTAL CONSIDERATIONS**

The leather industry is a chemical processing industry. Processing is largely carried out in a water medium and as such, there is potential for a number of environmental concerns.
The regulatory framework in New Zealand under the Resource Management Act is quite strong and in essence is set up to ensure that resources are used in such a way that their use is not harmful to the environment. The framework governs discharges to air, water and on land and the responsibility is to ensure that these are not adversely affected.

The main tanneries in New Zealand are sited in areas where their waste waters can be discharged into metropolitan sewerage systems and treated along with domestic sewage. However, as the metropolitan authorities are also governed by the Resource Management Act, standards are set which must be met before discharge to the sewers can take place. These vary from place to place depending on local conditions, but are set so that the ultimate discharge to natural receiving water reaches acceptable standards.

It can be seen from the flow chart (Figure 2) that the main waste products originating from leather processing are hair or wool, salts, alkali (lime), sulphides, acids, chromium salts and dye-stuffs. Hair and wool are useful by-products, the hair being used for felt manufacture and wool for clothing and carpeting. Salt content presents no major difficulty while the acids and alkalis can be used to neutralise one another. Any excess of acid or alkali can be readily and cheaply neutralised.

The two components of normal tannery effluent which are difficult to deal with are sulphides and chromium salts. Each is undesirable for different reasons.

**Sulphide**

Solutions of sulphides in neutral or acidic conditions liberate significant quantities of hydrogen sulphide, which has a strong and objectionable smell (like rotten eggs) and is quite toxic - a potential hazard for sewerage workers. The main objection to sulphide, however, is its corrosive action on the sewer fabric. Hydrogen sulphide, liberated in acidic or near-acidic conditions, dissolves in condensed moisture in the sewers above the water line. This is oxidised by various bacteria into sulphuric acid. The bacteria require water, oxygen, carbon dioxide, ammonia and a source of oxidisable sulphur to reproduce. The pH must be below 7.5. These conditions are readily obtained due to the equilibrium:

\[
\text{CO}_3^{2-} \rightarrow \text{HCO}_3^- \rightarrow \text{CO}_2
\]

The pH under these conditions will soon get down to below 7.5 and eventually, by the activities of the bacteria, concentrations of up to 5% sulphuric acid can be built up. The sulphuric acid erodes the sewer walls by forming sulphates of the calcium, aluminium and iron found in concrete. Besides being readily soluble, these salts expand as they are formed and cause crumbling of the aggregates in the concrete.

The concentration of sulphide in solution can be considerably reduced by oxidation of the sulphide to some unoxidisable form of sulphur. This can be achieved by aeration of the industrial effluent by means of air injection in the presence of manganese ions as a catalyst. Under these conditions the sulphide content can be reduced to 2-3 mg per litre, which is acceptable to drainage authorities.
Chromium salts are undesirable for two reasons. Firstly, they can have a poisoning effect on the micro-organisms used for sewage treatment. Secondly they are precipitated as chromium hydroxide at neutral and near neutral pH's, meaning that they persist into the sludge from the sewage treatment plants. This gives a build-up of chromium (III) hydroxide in the sludge, which is undesirable as sludge is commonly used as a fertilizer. However, chromium is an essential trace element and there is no evidence of leaching of chromium in any form from sludges under normal conditions. As recently as 1994, the American courts prohibited the setting of limits on chromium(III) discharges as no detrimental effects had been demonstrated.

However, in most parts of the world there are still limits imposed.

Tannery effluents are discharged in an alkaline condition. Under these circumstances any chromium present is in the form of chromium hydroxide, a light, finely divided precipitate, which is very difficult to remove. Treatment consists of the addition of more alkali and trace quantities of polyelectrolyte flocculating agents. This aggregates the finely divided particles and greatly improves the rate of sedimentation so the chromium hydroxide can be removed in a normal settling operation, or by dissolved air flotation.

However, because of the limited world supply of chromium, it is necessary to develop means by which chromium tanning materials are recycled. This can be achieved in the main by saving and re-using the chromium tanning liquors. What tanning liquor is lost can be made alkaline and the chromium hydroxide separated and recovered by re-acidification.

The chromium sulphate so produced is suitable as a tanning solution.

Leather production requires the use of large quantities of water. Through the use of modern equipment efforts are being made to reduce this volume. By use of the methods described above, the levels of undesirable chemicals in the water discharged from the tanneries can be reduced to levels which are acceptable to drainage authorities and if necessary acceptable for discharge into surface waters. As with all pollution control, the costs are high, and significant volumes are involved. However, the technology now exists for these problems to be controlled.

CONCLUSION

The first leather was produced in pre-historic times. It was discovered quite by accident that a skin which has been soaked for some time in water which had been infused with the bark of trees did not rot or dry up like other skins, but remained soft and lasted for a long time. The
new material had many applications: it was useful for clothing, foot protection, for tying and binding and for tool making.

This discovery of the secret of making leather, like that of making fire, was one of the more important in early human history. From the initial accidental discovery until quite recent times, leather-making has remained a craft industry. It has developed through trial and error, with the scientific reasons behind the results being somewhat obscure. It can be seen that the chemistry involved is quite complex, and, as we are dealing with a chemically complex raw material, this will remain the situation for some time yet. However, as we obtain a better understanding of the chemistry of collagen, keratin and the various tanning materials used, the results obtained are becoming more predictable and more controllable.

Leather-making is now a scientifically based industry, but still retains some of the charm and mystery of the original craft. As such, it holds a constant challenge for the chemist who, through research and an understanding of the underlying principles, can control the quality of the final product to a much greater degree.

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