Barley for Brewing: Characteristic Changes during Malting, Brewing and Applications of its By-Products

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Barley for Brewing: Characteristic Changes during Malting, Brewing and Applications of its By-Products

Mahesh Gupta, Nissreen Abu-Ghannam, and Eimear Gallaghan

ABSTRACT: Barley is the basic raw material for brewing. Its chemical composition, brewing, and technological indices are highly determinative for the beer quality and the economical efficiency of the brewing process. Barley is rich in protein, carbohydrates, dietary fibers, minerals, and vitamins. The presence of nonstarch polysaccharides as mixed linkage (1-3), (1-4)-β-D-glucans and arabinoxylans together with the enzymes are responsible for barley modification. Malting is a complex process that involves many enzymes; important ones are α-amylase, β-amylase, α-glucosidase, and limit dextrinase. During the process of malting and brewing, the by-products left after separation of the wort are rich in protein, fibers, arabinoxylans, and β-glucan. This review summarizes and integrates barley grain with respect to nutritional, functional, and compositional changes that take place during malting and brewing. It also explores in-depth the several by-products obtained after brewing and their potential for various food applications. Barley brewing by-products offer an opportunity for cereal-based baked and extruded products with acceptable sensory and nutritional characteristics.

Importance of Barley Grain

Barley (Hordeum vulgare, vulgare L.) is a highly adaptable cereal grain that is produced in climates ranging from sub-Arctic to subtropical. It ranks 5th among all crops in dry matter production in the world today (129 million metric tons, 2002 to 2005 mean). Historically, barley has been an important food source in many parts of the world, including the Middle East, North Africa, and northern and eastern Europe (mainly Iran, Morocco, Ethiopia, Finland, England, Germany, Denmark, Russia, and Poland), and in Asia (Japan, India, Tibet, and Korea) (Chatterjee and Abrol 1977; Newman and Newman 2006). At present, only 2% of barley is used for human food (Baik and Ullrich 2008).

Barley grain is an excellent source of soluble and insoluble dietary fiber (DF) and other bioactive constituents, such as vitamin E (including toco-tri-enols), B-complex vitamins, minerals, and phenolic compounds. β-Glucans, the major fiber constituents of barley, have been implicated in lowering plasma cholesterol, improving lipid metabolism, and reducing glycemic index. The effectiveness of barley β-glucans in food products for lowering blood cholesterol has been documented in a number of studies (Newman and others 1989; Behall and others 2004). Barley is a rich source of tocols, including tocopherols and tocotrienols, which are known to reduce serum low-density lipoprotein cholesterol through their antioxidant action (Qureshi and others 1986). Whole grains are known for their fiber content, and therefore lower energy density, and as a source of vitamins and mineral components, both of which may increase satiety and reduce energy intake (Slavin 2003). In Western countries, pearled barley, whole, flaked, or ground is used in breakfast cereals, stews, soups, porridge, bakery flour blends, and baby foods. In Middle Eastern and North African countries, barley is pearled and ground, and used in soups, flat bread, and porridge (Bhatty 1993). Newman
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Barley grain that is clean, bright yellow-white, plump, thin-hulled, medium-hard, and uniform in size is generally suitable for food uses and preferred for pearling (Pomeranz 1974). Grain hardness is an important characteristic of barley because it determines the pearling and subsequent end-use quality of barley. Malt barley varieties are usually soft, whereas nonmalt barley varieties are usually hard. Psota and others (2007) also reported significant relationships between hardness of barley grain as assessed using the particle size index and hot water extract of malt as well as the malt quality index of barley malt. Other structural and compositional characteristics of barley endosperm could contribute to grain hardness, including proteins, starch, β-glucan, and their interactions, and packing during grain filling (Henry 1988). Generally, sound barley grain has a bright light-yellow or off-white color. Discolored barley grain often develops undesirable flavors when malted and has poor germination energy and vigor (Li and others 2003). The grain color of barley can vary from light yellow to purple, violet, blue, and black, which is mainly caused by the level of anthocyanins in the hull, pericarp, and/or aleurone layer. Highly colored types are also receiving attention for applications in functional foods due to their antioxidant properties (Satute-Gracia and others 1997; Nam and others 2006; Philpott and others 2006). However, most of the barley that is produced possesses bright, light yellow grain color, which is generally preferred for malting, brewing, and food purposes.

Whole barley grain consists of about 65% to 68% starch, 10% to 17% protein, 4% to 9% β-glucan, 2% to 3% free lipids, and 1.5% to 2.5% minerals (Czuchajowska and others 1998; Izydorczyk and others 2000; Quinde and others 2004). Total DF ranges from 11% to 34% and soluble DF from 3% to 28% (Fastnacht 2001). Pearl barley reduces the contents of insoluble fiber, protein, ash, and free lipids (Quinde and others 2004). On the other hand, hulled barley is preferred to hull-less barley for malting and brewing because of the contribution of the hull to beer flavor and as a filtering aid during brewing (Burger and La-Berge 1985). The amylose content of barley starch varies from 0% to 5% in waxy, 20% to 30% in normal, and up to 45% in high-amylose barley (Bhatty and Rossnagel 1997). β-glucans constitute approximately 75% of the barley endosperm cell walls together with 20% arabinoxylans and protein. Both β-glucans and arabinoxylans determine wort viscosity and beer filtration rates (Stewart and others 2000), and form a barrier for hydrolytic enzymes attacking starch and protein within the cell walls causing potential health benefits such as prevention of constipation, reduction in risk of colorectal cancer (Bingham 1990; Faivre and Bonithon-Kopp 1999), lowering of blood cholesterol, and controlling diabetes management (Gallagher and others 1993; Frost and others 1999). Barley endosperm protein is rich in prolamin storage proteins (hordeins) and has moderate nutritional quality (Newman and McGuire 1985). High-lysine barley mutants, which contain 2% to 3% greater lysine than normal lysine types could provide high-quality, protein-enriched barley grains for the human diet (high lysine content of 5% to 6% compared to 3% as normal ones) (Ulrich and Eslick 1978).

A large number of parameters have been proposed to define malting quality. It is also a fact that the texture of the endosperm influences the malt modification process by affecting water uptake and consequently enzyme synthesis and movement within the endosperm (Chandra and others 1999). Anderson and others (1999) studied the variation and correlation between chemical and physical characteristics of barley samples including kernel hardness, but found only a low correlation between kernel hardness and physical and chemical grain properties. One another factor as potential influence of sulfur (S) on barley malting quality has so far received little attention. Sulfur deficiency has been shown to affect the composition of proteins in barley grain, with depletion in the S-rich B hordein and the high-molecular weight (HMW) D hordein and an increase in the S-poor C hordein (Shewry 1993). The melting of hull-less barley, however, presents a number of challenges due to differences in chemical and physical characteristics.

Characteristics of Malting Barley Grain

Barley is the primary cereal used in the production of malt in the world. Two types of barley are frequently used for the malthoring process: 6- and 2-row. Two-row barley produces malt with a large extract, lighter color, and less enzyme content than the 6-row type (Broderick 1977). From the different quality parameters reported in the literature, hot-water extract (HWE), kernel size fractions, kernel weight, β-glucan and protein contents, malting losses, friability, α-amylase activity, viscosity, and soluble nitrogen ratio (SNR) are common assays used to test the quality of barley malt (Fox and others 2003). In addition, fast hydration and germination are necessary traits of barley for good malting quality (Ulonska and Baumer 1976; Briggs 1998). During mashing, barley undergoes an incomplete natural germination process that involves a series of enzyme degradations of barley kernel endosperm. As a result, this enzyme degradation, endosperm cell walls are degraded, and starch granules are released from the matrix of the endosperm in which they are embedded. These structural changes and biochemical degradations of the endosperm components are referred to as endosperm modification (Gunkel and others 2002). Maling is defined as the controlled germination of cereals, to ensure a given physical and biochemical change within the grain, which is then stabilized by grain drying. Three process steps are necessary to ensure that these changes occur: (1) steeping, to ensure good absorption of water by the grain (from 12% to at least 40% of moisture); (2) germination, to maintain embryo growth, enzyme synthesis and a limited endosperm breakdown; and (3) kilning, to ensure product stability.
Different kernel properties have been identified as factors affecting water uptake during steeping of barley, for example, endosperm structure, starch content, protein content, and cell wall properties (Ogushi and others 2002). Loosely packed endosperm gives soft (mealy) structure and facilitates better moisture and enzyme movements in the endosperm. Thus, a mealy endosperm is more easily degraded by hydrolytic enzymes during malting (Swanson and others 1995). On the other hand, starch granule size and distribution, amylose, amylopectin, β-glucan, and arabinoxylan content have also been proposed as factors in affecting the hardness of the endosperm (Dombrink and Knutson 1997; Tohno-Oka and others 2004). As a result of the malting process, there is an increase in enzyme activity, soluble protein, and breakdown of starch into simple sugars, along with development of the typical color and flavor (Hoseney 1994). The final moisture content of malt is approximately 35 to 40 g/kg, being a highly hygroscopic product. Mashing is a key step in the beer production process. During mashing, enzymatic degradation of the polysaccharides present in the malt takes place. Fermentable carbohydrates are produced from the degradation of the polysaccharide starch. Such carbohydrates are converted into alcohol in the fermentation step of the beer manufacturing. Nonstarch polysaccharides also degrade during mashing into smaller chain carbohydrates. Different enzymes catalyze all the involved reactions. Because the activity of the different enzymes is highly dependent on temperature, the manipulation of such variable is the main control mechanism for the mashing process (Hardwick 1995).

**Various Factors Affecting Brewing Process of Barley Grain**

**Properties of antioxidants**

Antioxidants are not evenly distributed in barley grains. Salomonsen and others (1980) indicated that p-coumaric acid was present in the lowest amount in the center of the barley kernel and rapidly increased toward the outer layers, such as lignified husk (Maillard and Berset 1995), whereas Goupy and others (1999) have indicated that phenolic acids were mainly present in the aleurone layer and endosperm. The content of ferulic acid is highest in the cell walls of the aleurone layer, which is rich in arabinoxylans. Maillard and Berzet (1995) have suggested that trans-ferulic acid, trans-p-coumaric acid, and cis-ferulic acid from barley and malt. The natural antioxidants in cereals may act as free radical scavengers, reducing agents, potential complexes of pro-oxidant metals, and singlet oxygen quenchers (Zielinski 2002). Moreover, many of the natural antioxidants present in barley exhibit a wide range of biological effects, including antibacterial, antiviral, anti-inflammatory, anti-allergic, and antithrombotic effects, and may also be involved in vasodilatory actions (Cook and Sammon 1996). Polyphenols identified in barley include anthocyanins, flavonols, phenolic acids, catechins, and proanthocyanidins (Goupy and others 1999). There are more than 50 proanthocyanidins reported in barley, and they include oligomeric and polymeric flavan-3-ol, catechin (c), and galloカテchin (gc). The most abundant proanthocyanidins in barley are dimeric proanthocyanin B3 and proanthocyanin B3. Major trimers include T1 (gc–gc–c), T2 (gc–c–c), T3 (c–gc–c), and T4 or proanthocyanidins C2 (c–c–c) (Friedrich and others 2000). Phenolic compounds in cereal grains exist in the free, soluble esters or conjugates, and insoluble-bound forms (Adom and Liu 2002). Antioxidants are generally thought to play a significant role in mashing and brewing due to their ability to delay or prevent oxidation reactions and oxygen free radical reactions. Antioxidants such as sulfites, formaldehyde, or ascorbate, can be added into the brewing process to improve beer flavor stability. About 80% of phenolic compounds present in beer are derived from barley malt, and the remaining come from hops (Goupy and others 1999). Those phenolic compounds in malting barley include polyphenols (benzolic and cinnamic acid derivatives), flavonoids, proanthocyanidins, tannins, and amino phenolic compounds (Hernanz and others 2001; Bonoli and others 2004), all of which are known to inhibit nonenzymatic lipid peroxidation and widely recognized as having important antioxidant and antiradical properties. Therefore, the presence of the natural antioxidants in malting barley and screening of malting barley variety with the highest level of radical scavengers seems very important to produce beers with high levels of antioxidant activity (Maillard and others 1996).

**Arabinoxylans and β-glucan content**

Arabinoxylans (AX) consist of a linear-chain backbone of β-D-xylpyranosyl (Xylp) residues linked through (1-4)-glycosidic linkages. α-L-Arabinofuranosyl (Ar) residues are attached to some of the Xylp residues at O-3, O-4, and/or at both O-2, 3 positions, resulting in 4 structural elements in the molecular structure of arabinoxylans: monosubstituted Xylp at O-2 or O-3, disubstituted Xylp at O-2, 3, and unsubstituted Xylp (Figure 1; Gruppen and others 1993; Izydorczyk and Biliaderis 1995; Vink and others 1999). A unique feature of arabinoxylans is the presence of hydroxycinnamic acids and ferulic and p-coumaric acids, esterified to O-5 of Aral linked to O-3 of the xylose residues (Smith and Hartley 1983). The content of arabinoxylans in barley also depends on genetic and environmental factors (Fleury and others 1997; Izydorczyk and others 2000; Holtekjølen and others 2007) but appears to be less variable than that of β-glucans. Compared to other grains, the amount of arabinoxylans in barley is similar to that in wheat (5.8%), but higher than in oats (2.7% to 3.5%), sorghum (1.8%), or rice (2.6%) (Izydorczyk and Biliaderis 2007). Fleury and others (1997) reported that the amount of arabinoxylans in the hull-less barley (3.37% to 4.30%) was significantly lower than in the 2- or 6-rowed (5.41% to 6.42%) covered barley, and linked the differences to the absence of hulls in the former. Six-rowed barley cultivars generally contain slightly higher levels of arabinoxylans than 2-rowed cultivars (Fleury and others 1997). The presence of the waxy gene in barley does not affect the content of arabinoxylans to the same extent as that of β-glucans. In view of the importance of nonstarch polysaccharides in the malting of barley and in subsequent steps in the brewing process, including the possible influence of residual cell wall on the rate of wort separation (Han and Schwarz 1996). But undermodified malts, the fact that arabinoxylans cannot be degraded sufficiently, may cause many problems such as low extract yield, high wort viscosity, decrease of filtration rate, and haze formation in brewing (Coote and Kirsop 1976). It has been found that other large molecules such as AX, proteins, and polyphenols were also associated with reduced beer filtration, in particular microfiltration. In fact, it has been reported that the amount of AX in commercial beer is approximately 10 times greater than that of β-glucan (Schwarz and Han 1995). The enzymes that degrade AX are often produced late in the germination process (Banik and others 1997), and high levels of AX can survive through the brewing into the final beer. Some AX are solubilized from the cell walls but are not extensively degraded by endogenous enzymes during malting (Voragen and others 1987). Malt extracts can contain high levels of AX and cause difficulties associated with the filtration of viscous extracts may significantly deteriorate the performance of the brewing processes (Bamforth 1985).

Mixed linkage (1-3,1-4)-β-D-glucans, commonly known as β-glucans, are linear homopolymers of D-glucopyranosyl (GlcP) residues linked mostly via 2 or 3 consecutive β-(1-4) linkages
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Figure 1 — Structural elements present in arabinoxylans: (A) unsubstituted Xylp, (B) monosubstituted Xylp at O-2, (C) monosubstituted Xylp at O-3 with ferulic acid residue esterified, and (D) disubstituted Xylp at O-2,3 (from Izydorczyk and Dexter 2008).

Figure 2 — General molecular structure of \( \beta \)-glucans and hydrolysis products obtained upon digestion of \( \beta \)-glucans with lichenase (from Izydorczyk and Dexter 2008).

(Figure 2) that are separated by a single \( \beta \)-(1-3) linkage (Cui and others 2000). The changes of both (1-3, 1-4)-\( \beta \)-D-glucan (referred to as \( \beta \)-glucan) content and (1-3, 1-4)-\( \beta \)-D-glucan-4-glucanohydrolase (referred to as \( \beta \)-glucanase, E.C.3.2.1.73) activity in barley during malting are important for malt producers, which is closely associated with malt yield and quality. \( \beta \)-Glucanase is a cell wall polysaccharide, which accounts for approximately 70% (w/w) of the endosperm cell in barley (Forrest and Wainwright 1977; Jeraci and Lewis 1989). Compared to other grains, barley contains a relatively high concentration of \( \beta \)-glucan, a viscous and fermentable DF, therefore may be highly satiating (Marciani and others 2001). In the brewing industry, a high content of \( \beta \)-glucan in barley may lead to insufficient degradation of cell walls, which in turn hinders the diffusion of enzymes, germination, and the mobilization of kernel reserves, and hence reduces malt extract. Residual \( \beta \)-glucan may also lead to highly viscous wort, giving rise to a filtration problem in the brewery, and it may participate in maturing of beer, causing chill haze (Bamforth 1982). The degradation of endosperm cell walls and subsequent changes in \( \beta \)-glucan levels during malting are, to a great extent, related to \( \beta \)-glucanase activity, which depolymerizes \( \beta \)-glucan (Etokakpan 1993). Therefore, better malting performance is expected to be associated with lower levels of \( \beta \)-glucan in grains and higher levels of \( \beta \)-glucanase in malt. Historically, reduced beer filtration efficiency has been mainly attributed to \( \beta \)-glucan in the brewing process. \( \beta \)-Glucan may increase the viscosity of beer by forming gels, consisting primarily of HMW \( \beta \)-glucan molecules (Home and others 1999). Barley has gained popularity due to the functional properties most likely antioxidant and radical scavenging activity due to its bioactive compounds such as \( \beta \)-glucan, arabinoxylan, oligosaccharides, tocols, and phenolic compounds (Baik and Ullrich 2008).

Protein in barley grain

Proteins are among barley components that are essential for the quality of malt and beer. First, high-protein contents decrease available carbohydrates, with a negative influence on the brewing process (Peltonen and others 1994; Fox and others 2002) and second, proteolysis (protease hydrolysis producing amino acids and peptides from hordeins) during malting and mashing is necessary for yeast metabolism (Moll 1979). Finally, soluble proteins are important in beer head retention and stability. The
protein content in barley grains represents, approximately, 8% to 15% of its total mass. Hordeins are the most abundant proteins (40% to 50%) found in a barley grain (Osman and others 2002). In addition to the hordeins, other proteins have been identified, including albumins, glutelins (globulins), friabilin, enzymes, and serpins (Finnie and others 2002; Fox and others 2002; Osman and others 2003; Boren and others 2004). Barley hordeins are divided into 5 groups based on their electrophoretic mobilities and amino acid compositions: the B and C hordeins (70% to 80%) and 10% to 20% of the hordein fraction, respectively) and D, Z, and γ hordeins (less than 5% of the total hordein fraction) (Shewry 1993; Tatham and Shewry 1995). The B hordeins can be subdivided into B1, B2, and B3 subtypes (Skerritt and Janes 1992). Furthermore, a distinction is made between the sulfur-rich (B and γ hordeins), the sulfur-poor (C hordeins), and HMW prolamins (D hordeins) (Shewry 1993; Tatham and Shewry 1995). It has been hypothesized by Moonen and others (1987) that in barley, HMW subunits form a backbone, which binds low-molecular-weight (LMW) subunits through disulfide bridges to form a gel-like aggregate. The majority of beer protein lies in the 10 to 40 kDa size range (Leiper and others 2003). Mostly, the origin of HMW protein is malted barley (Hughes and Baxter 2001). Some beer proteins appear to have no function in beer except their contribution to mouthfeel, flavor, texture, body, color, and nutritional value (Leiper and others 2003; Osman and others 2003). Protein Z, LTP1 (lipid transfer protein), and other proteins present in beer have been associated with foam formation and/or stabilization (Evans and Sheehan 2002; Perrocheau and others 2005). Protein Z has also been related to beer haze (Curioni and others 1995). During malting, barley proteins are in part degraded to amino acids and small peptides by a range of proteolytic enzymes (Baxter 1982; Bax and others 1986; Jones 2005a, 2005b). Brewer's spent grain (BSG), the main by-product of the brewing industry, is rich in proteins and DF (Musatto and others 2006). Identification of protein in the malt and beer samples by polyacrylamide gel electrophoresis or high-performance liquid chromatography has become a routine laboratory test in grain segregation in malt houses and in barley breeding programs.

Hydrolysis of starch

Rapid hydrolysis of starch to the fermentable carbohydrates glucose, maltose, and maltotriose is an important aspect of brewing. Starch hydrolysis is carried out by the malt enzymes α-amylase, β-amylase, limit dextrinase, and α-glucosidase (Manners 1985). Limit dextrinase is responsible for hydrolyzing the (1→6)-α-glucosidic branch points in LMW branched dextrins formed by the action of α- and β-amylase on starch components (Manners and others 1970). Starch granules can be encapsulated by a rigid protein matrix or by cell walls (Weurding and others 2001). α-Amylase can solubilize both amorphous and crystalline regions (Lauro and others 1993) of starch granules attacking the (α-4)-linkages of starch producing oligosaccharides. β-Amylase also attacks (α-4)-linkages from the nonreducing ends of amylose and amylopectin molecules (Bamforth and Quain 1989; Lewis and Young 1995). A range of fermentable sugars is produced from the action of these enzymes on starch during the mashing process. These include glucose, sucrose, fructose, and mainly maltose and also some LMW dextrins (Slack and Wainwright 1980; Lauro and others 1993). Starch α-amylase is also active on the particle size of starch granules (Coloma and others 1988). Large starch granules gelatinize earlier than small ones at high temperatures, despite the fact that the small granules have a slightly lower gelatinization enthalpy and a higher surface-volume ratio than the large granules, and hence one would expect them to gelatinize earlier than the large ones (Soulaka and Morrison 1985; Morrison and others 1994). The rate of hydrolysis may be influenced by both the surface features and internal structure of starch granules (Li and others 2003). Fermented wort and beer, however, contains appreciable levels of branched dextrin (Enevoldsen and Schmidt 1973) and suggesting that there is limited hydrolysis by limit dextrinase during the mashing process. Results to date suggest that the enzyme is readily solubilized from malt, but most of it is in an inactive form that requires “activation” to release, in full, the enzymic activity (MacGregor and others 1994a, 1994b; Sisson 1996). Other enzymes in the gist, such as limit dextrinase, may also contribute to the fermentable sugar profile. These enzyme activities are profoundly influenced by such a high mashing temperature as 65 °C. β-Amylase, in particular, is rapidly denatured at temperatures above 55 °C. α-Amylase is rather more stable and remains active for over an hour at 65 °C (Muller 1991).

Role of Enzymes in Malting and Brewing of Barley Grain

The conversion of barley into beer represents mankind’s oldest and most complex example of applied enzymology. Indeed, historically some of the most significant advances in enzymology have been linked to the world of brewing, such as Eduard Buchner’s extraction of enzymes from brewing yeast (Buchner 1897) and Adrian Brown’s kinetic analysis of invertase (Brown 1902). In determining the factors that a bearing on the quality of beer, brewers have learned not only how the endogenous enzymes contribute to issues such as fermentability, filterability, foam, clarity, flavor, so on, but also how to take advantage of exogenous enzymes. There are 3 primary “enzyme reactor” stages in the conversion of barley to beer (Bamforth 2006): barley kernel, mash tun, and the yeast cell. Only in one of these, the mash tun, is considered a “typical” enzyme reactor and has been extensively researched (Boulton and Quain 2001).

More than 40 endoproteases have been identified in malt, broadly classified into cysteine-, metallo-, aspartic-, and serine-proteinas (Jones 2005a). There are also exo-peptidases classified into carboxypeptidases (Mikola and others 1971) and amino peptides (Sopanen and Mikola 1975). A substantive reason for the limited action of the endo-peptidases in mashing is the presence of inhibitor proteins (Jones 2005b). Principal among such inhibitors are lipid transfer proteins that block the cysteine-proteinas (Jones 2005b). Jones and Budde (2005) suggest that 32% of the soluble protein in malt is already in the ungerminated barley form, 46% is released in malting and the rest solubilized in mashing. It was shown that over the pH range 5 to 6.6, the proteolytic activity of malt can vary more than 7-fold (Jones and Budde 2003). Various factors may come together in causing the release of the enzyme in an active form during mashing (Bamforth and Briggs 2000). It was recently suggested that serine-proteinas have a key role to play here (Schmitt and Marinac 2008). The least investigated of the endogenous starch-degrading enzymes in malted barley is α-glucosidase, although it has been claimed to be 2nd only to α-amylase for its importance in starch degradation during malting (Sun and Henson 1991). However, the enzyme is thermostable and likely to be of limited significance during mashing (Muslin and others 2000). Low-calorie beers, so-called Light, are the biggest selling style of beers and the exaggerated scares about “bad carbs” in beer (Bamforth 2005) led to the advent of “low carb beers.” For such products, glucoamylase and pullulanase are of great utility in dealing with the dextrins surviving because of the limited action of limit dextrinase (Goode and others 2005).

By-Products Obtained during Malting and Brewing

The brewing industry generates relatively large amounts of by-products and wastes spent grain, spent hops, and yeast being the most common. However, as most of these are agricultural
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products, they can be readily recycled and reused. Thus, compared to other industries, the brewing industry tends to be more environmentally friendly (Ishiwaki and others 2000). Spent grain is the most abundant brewing by-product, corresponding to approximately 85% of total by-products generated (Reinold 1997). According to Townsley (1979) spent grain accounts, on average, for 31% of the original malt weight, representing approximately 20 kg per 100 L of beer produced (Reinold 1997). BSG is available at low or no cost throughout the year and is produced in large quantities not only by large but also small breweries. In the brewery, malted barley is milled, mixed with water in the mash tun, and the temperature of mash slowly increased from 37 to 78 °C to promote enzymatic hydrolysis of malt constituents (Figure 3). This enzymatic conversion stage (mashing) produces a sweet liquid known as wort. The insoluble, undergraded part the malted barley grain is allowed to settle to form a bed in the mash tun and the sweet wort filtered through it (lautering) (Linko and others 1998; Dragone and others 2002). Figure 3 is a schematic representation of the process resulting in the production of brewers' spent grain from barley grain. BSG may consist of the residues from malted barley, or those from malted barley and adjuncts (nonmalt sources of fermentable sugars), such as wheat, rice, or maize added during mashing (Reinold 1997). The chemical composition of BSG varies according to barley variety, harvest time, malting and mashing conditions, and the quality and type of adjuncts added in the brewing process (Huige 1994; Santos and others 2003); but in general, BSG is considered as a lignocellulosic material rich in protein and fiber, which account for around 20% and 70% of its composition, respectively. Microscopic examination shows the presence of numerous fibrous tissues from the surface layers of the original barley grain (Figure 4). The main components of these fibrous tissues are arabinoxylan, lignin (a polyphenolic macromolecule), and cellulose (a linear homopolymer of glucose units). Analyses of BSG by Santos and others (2003) indicated that besides fiber, 24.2% protein, 3.9% lipid, and 3.4% ash are present in oven-dried BSG. The protein, apparent starch, nonstarch polysaccharide composition fraction is different in BSG from pilot scale trials of malting barley of different varieties. Protein and fiber are highly concentrated in spent grain because most of the barley starch is removed during mashing (Kissel and Prentice 1979).

Minerals, vitamins, and amino acids are also found in BSG. The mineral elements include calcium, cobalt, copper, iron, magnesium, manganese, phosphorus, potassium, selenium, sodium, and sulfur, all in concentrations lower than 0.5% each (Pomeranz and Dikeman 1976; Huige 1994). The vitamins include (in ppm): biotin (0.1), choline (1800), folic acid (0.2), niacin...
(44), pantothenic acid (8.5), riboflavin (1.5), thiamine (0.7), and pyridoxine (0.7); protein-bound amino acids include leucine, valine, alanine, serine, glycine, glutamic acid, and aspartic acid in the largest amounts, and tyrosine, proline, threonine, arginine, and lysine in smaller amounts. Cystine, histidine, isoleucine, methionine, phenylalanine, and tryptophan also present in minor quantity (Huige 1994).

**Novel Food Applications of the By-Products of Malting and Brewing of Barley**

BSG can be employed either as a wet residue, shortly after separation from the wort at lautering, or as a dried material (Townesly 1979; Ozturk and others 2002). According to Huige (1994), BSG is an excellent feed ingredient for ruminants because it can be combined with inexpensive nitrogen sources, such as urea, to provide all the essential amino acids. In addition to its high nutritional value, BSG is reported to promote increased milk production without affecting animal fertility (Sawadogo and others 1989; Belibasakis and Tsirgogianni 1996a, 1996b; Reinold 1997). When BSG was incorporated into the diet of cows, milk yield, milk total solid content, and milk fat yield were increased. At the same time, blood plasma concentrations of glucose, total protein, albumin, urea, triglycerides, cholesterol, phospholipids, sodium, potassium, calcium, phosphorus, and magnesium were not affected (Belibasakis and Tsirgogianni 1996a, 1996b). Kaur and Saxena (2004) evaluated BSG as a replacement for rice bran in a fish diet and observed that fish fed with a diet containing rice bran and 30% BSG had a superior body weight gain when compared with fish fed with rice bran only.

**Food applications**

BSG, the residue left after separation of the wort during the brewing process (Santos and others 2003), is rich in cellulose and noncellulosic polysaccharides, mainly arabinoxylans (Mandalari and others 2005) as well as protein and β-glucan (Mussatto and others 2006). Approximately, 3.4 million metric ton of spent grains from the brewing industry are produced in the European Union every year (Eurostat data 2005). These plant-derived waste co-products are known to contain significant amounts of valuable components, which remain unexploited waste in the current processes. Because of its high moisture and fermentable sugar content, BSG becomes an environmental problem after a short time (7–10 d). It has a strong potential for being recycled and used as a cheap source of fiber that may provide a number of benefits when incorporated into human diets such as for the prevention of certain diseases including cancer, gastrointestinal disorders, diabetics, and coronary heart disease (Aman and others 1994; Jacobs and others 1998). Because of its relatively low cost and high nutritive value, BSG has been evaluated for the manufacture of flakes, whole wheat bread, biscuits, and aperitif snacks. However, BSG is too granular for direct addition to food and must first be converted to flour (Hassona 1993; Miranda and others 1994a, 1994b; Ozturk and others 2002). A high-protein flour prepared from BSG was successfully incorporated into a number of bakery products, including breads, muffins, cookies, mixed grain cereals, fruit and vegetable loaves, cakes, waffles, pancakes, tortillas, snacks, doughnuts, and brownies (Townesly 1979; Huige 1994). Nevertheless, there are some limitations in the use of flour as a protein additive or as a partial replacement for presently used flours, due to its color and flavor. BSG is brownish in color when moist and thus can only be used in off-white products, such as light-colored cookies, cakes, bread, or spaghetti that are made entirely from whole meal flour. Moreover, because of alterations in the flavor and physical properties (for example, texture) of the final products, only relatively small quantities (5% and 10%) can be incorporated (Townesly 1979; Hassona 1993; Miranda and others 1994a, 1994b). Prentice and D’Appolonia (1977) made high fiber bread containing BSG and evaluated its consumer acceptance. The results of their study showed that incorporation of BSG increase the nutritional and consumer acceptance level. In another study, BSG was finely milled and heat-treated at 45, 100 or 150 °C and replaced white flour in a conventional bread formula at 5%, 10%, and 15% levels. Bread containing heat-treated (45 °C) BSG, at 5% and 10% flour replacement levels was accepted favorably (Hassona 1993). Some properties of BSG flour in foods are shown in Table 1. The ingestion of BSG or derived products provides benefits for health, and is associated with increased fecal weight, accelerated transit time, increased cholesterol and fat excretion, and decrease in gallstones (Fastnauht 2001). Incorporation of spent grain in rat diets prevented an increase in plasma total lipids as well as of cholesterol (Hassona 1993; Ishiwaki and others 2000). BSG has been converted to a new protein-rich fibrous foodstuff by separating the husk fraction by milling and sieving. The product, germinated barley foodstuff (GBF), contains the aleurone layer, scutellum, and germ fractions of germinated barley, and is composed mainly of noncellulosic polysaccharides and glutamine-rich protein and is low in lignin (Kanauchi and Agata 1997). GBF feeding is considered a potentially new attractive prebiotic treatment in patients with ulcerative colitis (Kanauchi and others 2001; Bamba and others 2002). Furthermore, GBF has a high water-holding capacity compared with other water-insoluble DF sources, and this feature might contribute to a conspicuously high stool-forming ability in the colon (improvement of bowel movement) (Bamba and others 2002). GBF also appears to be safe and well tolerated. On the whole, BSG is a cheap source of protein and fiber that may provide a number of benefits when incorporated in human diets. For this reason, it is a potentially important food ingredient, especially in developing countries where poor malnutrition exists.

<table>
<thead>
<tr>
<th>Table 1 — Properties of BSG flour in foods.</th>
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<tbody>
<tr>
<td>1. Ease of blending</td>
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<td>2. Calorie content is approximately half that of most cereal flours (27.0 MJ/kg)</td>
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<tr>
<td>3. High water absorption capacity</td>
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<tr>
<td>4. Provides valuable minerals such as Ca, P, Fe, Cu, Zn, and Mg</td>
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<tr>
<td>5. Low fat absorption (beneficial for batters and coating)</td>
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<tr>
<td>6. Uniform tan color, bland flavor, and mildly roasted aroma</td>
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<tr>
<td>7. High fiber content as arabinoxylans (21.8%)</td>
</tr>
<tr>
<td>8. High protein content (24%)</td>
</tr>
</tbody>
</table>

Data from Huige (1994).

aData from Okamoto and others (2002).

bData from Kanauchi and others (2001).
a cheap source of protein and fiber that may provide a number of benefits when incorporated in human diets (Mussatto and others 2006). However, reference searches indicate that very little attention on the incorporation of BSG into extruded products has been published. Extrusion is a continuous cooking and shaping (forming) process designed to give unique physical and chemical functionality to food materials. Raw food ingredients undergo many order–disorder transitions, such as breakdown of starch granules, protein denaturation, and complex formation between lipids and amylose during extrusion. Extrusion-processing also controls the water activity of ingredients. It is therefore useful in producing shelf-stable foods and, more important, in producing a variety of items like snack foods and breakfast cereals. BSG is rich in complex carbohydrates and protein so it can be used for the extrusion processing to make various snacks.

Other applications

BSG is rich in polysaccharides and also in associated proteins and minerals and thus is a substrate of high biotechnological value. In this respect, several possible applications of BSG in biotechnological processes have been evaluated. BSG has been successfully used as a substrate for cultivating species of Pleurotus, Agrocybe, and Lentinus (Schliefbach and others 1992). BSG had good biological efficiency and high nutritional value as a substrate for Pleurotus ostreatus, especially when water-rinsed BSG was used (Wang and others 2001). It has been proposed that BSG favors the growth of these mushrooms not only due to its high protein content (Townsey 1979), but also to its high moisture content and physical properties such as particle size, volume weight, specific density, porosity, and water holding capacity (Wang and others 2001).

Cereal brans with compositions and physical structures comparable with BSG have been used extensively as substrates for the production of commercial enzymes in so-called koji or solid-state fermentations (Chou and Rwan 1995; Aikat and Bhattacharyya 2000; Sangeetha and others 2004). For this reason, BSG has also been evaluated as an alternative substrate for enzyme production. BSG is an efficient substrate for xylanase production by a Streptomyces isolate from Brazilian cerrado soil (Nascimento and others 2002), and for the production of xylanase and feruloyl esterase by Streptomyces avermitilis (Bartolome and others 2002, 2003). α-Amylase production by Bacillus subtilis (Duvnjak and others 1983) and Bacillus licheniformis (Okita and others 1985) cultivated on BSG has also been reported.

The reuse of BSG in the brewing process could be attractive from the point of view of brewery economics. Roberts (1976) showed that a BSG extract (a spent grain pressing concentrate) was effective as an antifoaming agent in the fermenter; in addition, hop utilization was improved and the properties of the final beer were not affected when the BSG extract was added. Addition of untreated BSG to wort enhanced the fermentation performance of yeast (Kado and others 1999), but the flavor and taste of the resulting beer was not satisfactory. However, addition of a neutralized acid extract of BSG to wort enhanced yeast performance and produced beer of quality equal to that of beer fermented without spent grain. BSG sequentially pretreated with HCl and NaOH solutions has been evaluated as a carrier for immobilizing brewer’s yeast (Saccharomyces uvarum) (Branyik and others 2001, 2002, 2004a, 2004b).

The cell walls of the barley grain residues in BSG are rich in cellulose and noncellulosic polysaccharides, in particular arabinanoxylans, but also some residual (1-3, 1-4)-β-glucan. The cell wall polysaccharides can be degraded into their corresponding constituents by hydrolytic procedures (hydrothermal, enzymatic, or acidic). Upon hydrolysis, cellulose yields glucose and the noncellulosic polysaccharides xyllose, mannose, galactose, and arabinose, as well as acetic and hydroxycinnamic acids (Palmqvist and Hahn-Hagerdal 2000; Mussatto and Roberto 2004) and some of these products are of industrial significance as precursors of food-grade chemicals or as energy sources in microbial fermentations. Hydrothermal hydrolysis (autohydrolysis by acetic acid released from its esterified form on the arabinoxylans) treatment of BSG with water at 150 °C for 60 and 120 min gave a wide variety of arabinino-oligosaccharides with different structural features. The arabinino-furanosyl side-branches on the xylan backbone are readily hydrolyzed and are easily removed by this treatment. The higher thermal sensitivity of the arabinose components compared to xylose, leads to release of large amounts of free arabinose when the temperature of the process is increased; and to major amounts of xylo-oligosaccharides. Hydroxycinnamic acids (ferulic and p-coumaric acids) present in BSG have potential uses in the food industry (Bartolome and Gomez-Cordoves 1999; Bartolome and others 2003). Bartolome and others (1997) used an esterase from Aspergillus niger to release ferulic acid from BSG and observed that 3.3% of the total ferulic acid was released, but in the presence of a xylanase from Trichoderma viride increased the extraction up to 30% (Bartolome and others 2003).

Future Considerations

Barley is one of the most ancient crops, and it has evolved through domestication to today as a major world crop based on acreage and production. It has great potential to reclaim some of its prominence as a food grain, largely due to its high nutritional value. Starch is a unique component of barley grain that gives physical properties to food products, but barley also contains high contents of protein and β-glucan. However, because human consumption of barley and barley-containing food products has been insignificant as compared to other cereal grains, the development of new processes and food products has been neglected and there has been little effort to define quality requirements for food uses. The significance of β-glucan and tocols for human nutrition is well known, but little is known about the functional properties of β-glucan for making food products. Some of the traits preferred for specific food applications are known through investigations on incorporating barley into wheat-based food products. On the other hand, the functional properties of β-glucans in food processing and end-use quality, with the exception of malting and brewing are little known. Much of the more recent interest in the use of β-glucans in food systems has stemmed from their use as a functional DF. Innovative ways are being developed to bring DF into new appealing high-fiber products that contribute to the recommended DF intake (Natl. Academy of Sciences 2002). The development of new techniques to use this agro-industrial by-product is of great interest due to the large volumes of spent grain produced. More research is required to understand how the gel protein is disaggregated during malting and how the aggregates form during mashing.

Increasing efforts are being directed towards the re-use of agro-industrial by-products, from both economic and environmental standpoints. Brewer’s spent grain (BSG) is an abundant by-product that can be obtained from brewing companies worldwide. However, in spite of all the possible applications described, its use is still limited, being basically used as animal feed or simply as a land fill. BSG can be considered as an attractive adjunct for human food. BSG has been used, for example, to produce protein-enriched breads, which could be very useful in the poorer regions of the world where food is scarce. On the other hand, considering that carbohydrates are the major components, more attention should be paid to its conversion into soluble and fermentable sugars. Currently, a number of added-value bioproducts such as organic acids, amino acids, vitamins, and...
ethanol, butanediol, among others, are produced by fermentation using glucose or xylose as substrates. But there is little information about the residual barley proteins present in BSG and their interactions with other BSG polymers. A consequential benefit of the use of industrial by-products such as BSG is as raw materials in various food applications. In addition, from an environmental viewpoint, the elimination of industrial by-products represents a solution to disposal and pollution problems.

References
Barley brewing and by-product applications.


