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A Structured Approach to Recover Valuable Compounds from Agri-food Side Streams

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Abstract

Food side streams contain useful compounds such as proteins, sugars, polyphenols, and amino acids that might get discarded during processing. The concentration of these components may be low (e.g., fruit side streams are mainly composed by water, around 90%, while polyphenol content in rapeseed meal is less than 3% dry weight) and therefore effective separation techniques should be evaluated. The aim of this review is to identify the different process steps (like pretreatment, volume reduction, phase change, solid removal, purification, and formulation) required to recover high-value products from agri-food residues. It reviews different plant-based byproducts as sources (cereal bran, fruit pomace, oilseed meals, fruit wastewater) of valuable compounds and discusses the relevant technologies required for processing (such as extraction, adsorption, crystallization, drying, among others). A structured approach to design recovery processes presented focused on high purity products. This work demonstrates that multiple high-value products can be recovered from a single agri-food side stream depending on the processing steps and the origin source (strong and soft structures and wastewater).

Keywords Valorization · Food side streams · Protein · Polyphenols · Separation technologies

Introduction

The increasing waste generation and the limited availability of natural resources have motivated the scientific community to investigate possibilities to recover valuable products from different waste streams such as wastewater and agri-food residues.

Food industry is recognized as one of the most important industries in the world but generates a significant amount of waste (Virtanen et al., 2016). On December 2015, the European Commission established a package to motivate EU members to move in the direction of a more circular economy (European Commission, 2017) and has taken the issue of food waste generation with seriousness. According to European Commission (2014), most of the waste is produced by the food manufacturing sector, and by 2020 is expected to rise to 126 million tons compared to 96 million tons generated in 2007 (European Commission, 2011). In order to contribute to this initiative, side streams of food industry can become inputs of other processes as they are sources of proteins, lipids, complex carbohydrates, and nutraceuticals. Therefore, valorization and recovery of high-value products from food byproducts is an attractive area that has been investigated lately around the globe (Virtanen et al., 2016). Recently, it has been proven that the conversion of biomass waste to bulk chemicals might be more profitable than its conversion to animal feed or transportation fuel (Tuck et al., 2012). In addition, increasing protein demand due to population growth could be satisfied by using plant-based residues such as proteins from canola meal resulted to be competitive and suitable for human consumption (Wanasundara et al., 2016).

Among the generated food byproducts, plant-based byproducts include fruit pomace, oilseed meals, cereal brans, wastewaters, etc. These residues are rich sources or dietary fiber (cellulose, hemicellulose, and lignin) and could be used within the biorefinery concept to produce biofuels or biochemicals from their respective hydrolysates. However, proteins and other nutraceuticals, mainly phenolic compounds (presented in the outer layers of most plant-based products (Naczk & Shahidi, 2006)) could be recovered. Extraction of proteins and nutraceuticals might be a fundamental step in
order to recover more valuable compounds from food byproducts.

This review describes and discusses the different valuable products presented exclusively in plant-based byproducts, such as oilseed meals, pea pods, cereal brans, and fruit pomace, among others. This is followed by the discussion of the different process techniques needed to recover these valuable compounds. Commonly, in these agri-food side streams, valuable compounds concentration is low (diluted systems, e.g., wastewater), and additionally, the presence of other low values impurities such as off-flavors and insoluble solids represents a technological challenge. Recovery and purification of the valuable compounds presented will additionally depend on the composition and volume of the side streams. Therefore, it is important to define different process stages. Combining the information available in literature for processing (Bongers & Almeida-Rivera, 2009; Galanakis, 2015) with some modifications, a generalized process flow scheme as the one defined in Fig. 1, could be applied as a starting point for the process synthesis of agri-food side streams.

The objective of the pretreatment step is to break the strong-linked structure of (solid) agri-food residues, such as cereal bran, in order to facilitate following processing steps; this might lead to recovery and purification of dietary fibers, or to conversions on sugars into biofuels. It is important to keep in mind that not all agri-food residues possess this strong matrix; therefore, this step is dependent of the source and the products to be recovered. For voluminous fruit and vegetable wastewaters, a volume reduction step is suggested at the beginning of the purification train, with the objective to concentrate and decrease volumetric load in the following steps. The third step, phase change, aims to extract the valuable components to a (different) liquid phase. The remained solids are removed (solid removal) and could be additionally treated for recovery of other products such as dietary fiber. The obtained liquid stream undergoes to a purification step, where the different compounds are separated or converted into other (bio) chemicals (e.g., sugars into bioethanol) and finally the last stage corresponds to formulation. This last stage has a main objective to get the product to its final form, e.g., powders and emulsions. The different unit tasks that can be applied in each stage of the processing of agri-food products are additionally indicated in Fig. 1.

This work discusses the different technologies for processing agri-food side streams to recovery valuable compounds as described in Fig. 1. Moreover, it provides an overview of possible purification processes of proteins, polyphenols, and dietary fibers from different plant-based starting materials.

### Food Side Streams: Valuable Products

The co-streams from food can originate from several branches mainly divided in two main groups, plant- and animal-based. This study will only focus on plant-based byproducts and their valuable compounds. Plant-based byproducts can be additionally subdivided in four groups namely: (1) cereals, (2) roots and tubers, (3) oil crop and pulses, and (4) fruits and vegetables (Galanakis, 2012). The average composition and the distribution of the different nutrients from plant-based sources are presented in Fig. 2, while Table 1 summarizes different plant-based side stream sources and the valuable products that could be recovered.

Cereal grains possess three botanical parts, endosperm, germ and bran. The bran is usually separated from the cereal grain, during milling operations, as it might have negative effects with the final product (Rosa-Sibakov et al., 2015) such as darker colors. However, cereal brans are sources of...
Table 1 Sources of plant-based byproducts and the potential valuable compounds

| Source               | Side streams        | Valuable compounds                                                                 | Reference                                                                 |
|----------------------|---------------------|------------------------------------------------------------------------------------|---------------------------------------------------------------------------|
| **Cereals**          |                     |                                     |                                                                           |
| Barley               | Bran, Malt, Straw   | Polyphenols (hydroxycinnamic acids) Protein                                       | Ahmad et al. (2019); Szwajgier and Borowiec (2012)                        |
| Corn                 | Cods, Husk, Silks   | Dietary fiber (cellulose and xylans) Minerals (P, K, Mg)                           | Lau et al. (2019); Pfaltzgraff et al. (2013)                               |
| Oat                  | Bran, Straw         | β-glucan Polyphenols Proteins                                                     | Patsioura et al. (2011)                                                   |
| Rice                 | Bran, Straw         | Dietary fiber Polyphenols Lipids Xylans                                            | Prakash and Ramaswamy (1996); Sohail et al. (2017); Liu et al. (2019); Ahmad et al. (2019); Orthoefer (2005) |
| Wheat                | Bran, Straw         | Proteins Arabinoxylan β-glucan Polyphenols (ferulic, sinapic and p-coumaric, flavonoids) Cellulose Xylans | Balandrán-Quintana et al. (2015); Ahmad et al. (2019)                        |
| **Root and tubers**  |                     |                                     |                                                                           |
| Asparagus            | Roots               | Carbohydrates Phenolics (flavonoids and hydroxycinnamic acids) Protein Saponins Oil | Zhang et al. (2019)                                                        |
| Carrots              | Pomace, Peel        | Dietary fiber α- and β-carotene Sugars Uronic acids Carotenoids                     | Sharma et al. (2012); Nawirska and Kwaśniewska (2005)                     |
| Cassava              | Peels, Pomace, Bagasse | Starch Dietary fiber Polyphenols Lipids Starch                                      | Versino et al. (2015); Ubalua (2007); Mullen et al. (2015)                |
| Potato               | Peel, Stillage (distillery water), Pulp | Dietary fiber Proteins Polyphenols (chlorogenic and hydroxycinnamic acid) Sugars Vitamins (B) Amino acids Pectin | Camire et al. (1997); Arapoglou et al. (2010); Rodriguez de Soltillo et al. (1994); Mullen et al. (2015) |
| **Oil crops and pulses** |                     |                                     |                                                                           |
| Hazelnut/almond/peanut | Hard shells, Leaf, Hull | Phenolic compounds (catechin, hydroxycinnamic acids, phenol benzoic acid) | Siriwardhana and Shahidi (2002); Shahidi et al. (2007)                     |
| Oilseed              | Oilseed meals (rapeseed, sunflowers, flaxseed, cotton) | Proteins Polyphenols Dietary fiber | Lomascolo et al. (2012); Wanasundara (2011)                              |
| Pea                  | Pod, Husk, Broken grains, Powder, Cotyledon | Dietary fiber Proteins Phenolics (hydroxycinnamic acids) | Mateos-Aparicio et al. (2010); Mateos-Aparicio et al. (2012)              |
### Table 1 (continued)

| Source       | Side streams | Valuable compounds                                                                 | Reference                                                                                     |
|--------------|--------------|-------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|
| 
| Fruits and vegetables |             |                                                                                     |                                                                                                |
| Apple        | Peel         | Dietary fiber                                                                       | Sudha (2011); Mourtzinos and Goula (2019)                                                      |
|              | Pomace       | Pectin                                                                              |                                                                                                |
|              | Seeds        | Polyphenols (catechins and proanthocyanidins)                                        |                                                                                                |
| Pineapple    | Peel         | Dietary fiber                                                                       | Roda and Lambi (2019); Larrauri et al. (1997); Seguí Gil and Fito Maupoey (2018)             |
|              | Core         | Proteins (bromelain)                                                                |                                                                                                |
|              | Stem         | Starch                                                                              |                                                                                                |
|              | Shells       | Polyphenols (myricetin, salicylic acid, and tannic acid)                             |                                                                                                |
| Orange       | Peel         | Dietary fiber                                                                       | Espina et al. (2011); May (1990); Chedea et al. (2010); Aravantinos-Zafiris et al. (1992); Scordino et al. (2007) |
|              | Pomace       | Essential oils (limonene)                                                            |                                                                                                |
|              |              | Pectin                                                                              |                                                                                                |
|              |              | Phenolics (flavonoids)                                                               |                                                                                                |
|              |              | Carotenoids                                                                         |                                                                                                |
| Tomato       | Pomace       | Lycopene                                                                            | Lu et al. (2019)                                                                               |
|              | Peels        | Dietary fiber                                                                       |                                                                                                |
|              | Seeds        | Pectin                                                                              |                                                                                                |
|              |              | Polysaccharides                                                                     |                                                                                                |
|              |              | Proteins                                                                            |                                                                                                |
|              |              | Oil                                                                                 |                                                                                                |
| Broccoli     | Pomace       | Proteins                                                                            | Shi et al. (2019)                                                                              |
|              | Stems        | Dietary fiber                                                                       |                                                                                                |
|              | Leaves       | Polyphenols (chlorogenic, neochlorogenic, and quinic acids)                          |                                                                                                |

**Fig. 2** Plant-based sources composition. 

- **Cereals**: Bran (7%), Cellulose fibers, Phenolics, Minerals, Endosperm (80-85%), Starch, Germ (21%), Proteins, Minerals, Fat
- **Oilseeds**: Seed coat (10-12%), Cellulose fibers, Phenolics, Embryo (88-90%), Embryotic axis, Proteins, Oil, Polyphenols, Carbohydrates, Cotyledon, Protein bodies, Oil bodies, Carbohydrates
- **Pulses**: Hull (25%), Cellulose fibers, Phenolics, Kernel (75%) (cotyledons), Proteins, Starch, Carbohydrates
- **Fruits**: Epicarp (peel), Phenolics, Cellulose fibers, Mesocarp, Sugars, Carbohydrates, Proteins, Seed, Endocarp
- **Vegetables**: Blade, Phenolics, Cellulose fiber, Sugars, Proteins, Vitamins, Lateral veins, Stem or branches, Cellulose fibers
- **Roots & Tubers**: Skin, Phenolics, Cellulose fibers, Proteins, Cortex, Carbohydrates (starch, fibers), Medulla, Carbohydrates, Proteins, Minerals, Inner medulla, Carbohydrates

Adapted from Naczk and Shahidi (2006), Butnariu and Butu (2015), Carrillo-López and Yahia (2019), González-Pérez and Arellano (2009), Islam and Ma (2016), List (2016), Rosell and Garzon (2015), Shukla et al. (1992)
nutritional compounds such as polyphenols, dietary fibers, and minerals (Heiniö et al., 2016). Different studies have shown the potential of using wheat bran as a source of valuable compounds. Rosa-Sibakov et al. (2015) compared the application of wheat bran in different baking products to increase their nutritional value. The study of Ahmad et al. (2019) demonstrated the different polyphenols available in wheat, barley, millet, and sorghum brans, assessing the antioxidant profiles, showing that millet and sorghum brans contain higher antioxidant activities than wheat and barley. Other byproducts obtained from the cereal sources are husks and straw which are rich on dietary fibers, glucoarabinoxylans (Hollmann & Lindhauer, 2005), and proteins (Prakash & Ramaswamy, 1996). Wheat, oat, barley, and rice straw are rich lignocellulosic biomasses (Pfaltzgraff et al., 2013) and can be used in biorefineries to obtain second-generation bioethanol or building blocks for other chemicals.

Fruit and vegetable side streams involve peels, leaves, pomace, and kernels, which are generated depending of the processing technology (juices, jams canning, jellies, etc.). The side streams generated are mostly composed of water and hydrocarbons (80–90%) with a low percentage of fat and proteins (Mirabella et al., 2014; Mullen et al., 2015). Orange peel is a rich source of essential oils (limonene), carotenoids, phenolic antioxidants, and pectin (Aravantinos-Zafiris et al., 1992; Chedea et al., 2010; Espina et al., 2011; May, 1990). A broad range of food products can be derived from this fruit namely sweet orange oil, orange blossom, honey, or marmalade (Siles López et al., 2010). Residues from pigmented orange pulp were valorized by Scordino et al. (2007) and successfully treated to recover sugars, citric acid, and pectin. Moreover, citrus fruits are rich on flavonoids, D-limonene, pulp, molasses (sugars), and essences (Virtanen et al., 2016). Around 25% of the processed apple is represented by apple pomace (Rodríguez Madera et al., 2013) and it is a natural source for commercial pectin around the world. In addition, it is also characterized by a high content of carbohydrates (cellulose, hemicellulose) and important polyphenols such as catechins, flavanols, hydroxycinnamates, and anthocyanins (Mourtzinos & Goula, 2019). Grape pomace, the byproduct of wine production, are rich sources of dietary fiber, oil, and phenolic compounds (anthocyanins and flavanols) (Bordiga et al., 2019), while tomato pomace is rich source of lycopene (principal carotenoid) proteins, dietary fiber, and oil (Lu et al., 2019). In addition, olive byproducts, olive mill wastewater, pomace, leaves, and seed are important sources of phenolic compounds, pectin, polysaccharides, and lignocellulosic fibers (Nunes et al., 2016).

According to the Food and Agriculture Organization of the United Nations (FAO) (2020), cassava, potatoes, sweet potatoes, yams, and carrots are the main root and tubers produced worldwide. Many bioactive compounds can be found in the byproducts of these food products, such as dietary fiber, proteins, antioxidants, and starch (Arapoglou et al., 2010; Camire et al., 1997; Mullen et al., 2015; Nawirska & Kwaśniewska, 2005; Rodriguez de Soltillo et al., 1994; Sharma et al., 2012; Ubalua, 2007; Versino et al., 2015). For instance, potato peel composition consists mainly of dietary fiber (around 50%)
Pulses (Mullen et al., 2015) and phenolic compounds (Schieber & Saldaña, 2009). Polyphenol antioxidative activity, from potato peel extracts, has been evaluated by Rodríguez de Soltillo et al. (1994) showing a similar performance of butylated hydroxyanisole (BHA) which is a food additive commonly used in food products to prevent rancidity (Prival, 2003). Carrot pomace accounts around 50% of the raw material during carrot juice production. This pomace still contains a significant amount of α- and β-carotene, which can be recovered and used as functional ingredient (Stoll et al., 2003). Similarly, sweet potato peels contain important antioxidants with chlorogenic acid, the highest phenolic in the root tissues (Truong et al., 2007).

At last, oilseed (flaxseed, canola/rapeseed, sunflower, and cotton) meals are rich sources of proteins, mainly storage proteins of two types, globulins and albumins (Lomascolo et al., 2012; Wanasundara, 2011). Fleddermann et al. (2013) evaluated the amino acids composition of canola meal proteins and compared with the one from soy protein for human nutrition. The authors demonstrated that the bioavailability of the proteins from canola protein isolate and soy protein isolate is similar, proving that canola proteins are relevant for nutrition (Fleddermann et al., 2013). In addition, oilseed meals are sources of polyphenols, such as sinapic acid (mayor phenolic acid in rapeseed) (Vuorela et al., 2004) which possesses antioxidant, antimicrobial, and anti-inflammatory properties (Nićiforović & Abramović, 2014).

Beans, chick peas, lentils lupins, and peas are the most cultivated and consume pulses (Food and Agriculture Organization of the United Nations (FAO), 2020). The byproducts generated after processing pulses include broken grains, husks, powder, unprocessed seeds, and shriveled pulses (Mullen et al., 2015). Mateos-Aparicio et al. (2010) evaluated byproducts of pea, broad bean, and okara (soybean byproduct) as rich sources of dietary fiber and polyphenols (Mateos-Aparicio et al., 2012). Moreover, the authors identified high quantity of vegetable proteins (around 30% dry matter) and fat (8.5% composed by linoleic and oleic acid) in okara which could be potentially recovered.

Using the process scheme presented in Fig. 1 and applying well-established methodologies such as the 5-stage universal recovery strategy proposed by Galanakis (2012) or the product-driven process synthesis (Almeida-Rivera et al., 2016), the processing of these agri-food side streams could be defined.

**Plant-Based Byproducts Processing: Recovery and Purification Techniques**

After the identification of the valuable components, processing of food side streams will undergo several steps (Fig. 1) and actual process synthesis will require different unit tasks. Depending on the source origin, some of the steps presented in Fig. 1 could be removed. In this work, source origin is divided in three groups: (1) lignocellulosic biomass which is characterized by plant fibers with a strong structure (e.g., cereal brans, oilseed meals), (2) fruit and vegetable pomaces (soft structures), and (3) fruits and vegetable wastewater (e.g., olive mill wastewater) (Fig. 3).

Considering strong structure side streams, such as bran from cereals, it has been proved that mechanical and thermal treatments (Agbor et al., 2011) of these solids improve the subsequent steps of the processing (pretreatment). The next step would correspond to the extraction of the components from the plant-based matrix to a liquid state (phase change). Depending on the conditions and medium to be used during the extraction, other components (valuable and non-valuable) might be co-extracted; therefore, separation and purification of the target components is required (solid removal and purification). Lastly, the final product will be formulated with a drying phase to remove water and generate the product form (e.g., powder).

For soft matrixes, such as fruit pomaces, pretreatment might not be needed and extraction of valuable products could be directly applied followed by purification and formulation, while for food wastewater streams, a volume reduction step is important in order to concentrate the streams and reduce the large volume generated. This will additionally benefit the overall process as smaller equipment would be needed for processing.

The following paragraphs of this section describe and discuss the different technologies that have been applied for processing plant-based residues (milling and pretreatment, extraction, purification, and formulation). It gives especially attention to adsorption as it proves to be the most promising purification technique for recovery of valuable products. Note that the downstream processing of the plant-based side streams will be dictated by the components to be recovered and the nature of the source.

**Milling and Pretreatment of Agri-food Byproducts**

Milling is a common operation in cereal, legumes, and oilseed processing as this operation is generating several byproducts such as hulls, husk, seed coat, and bran, among others. As previously mentioned, these byproducts are rich in bioactive compounds such as polyphenols, proteins, or dietary fibers (cellulose, hemicellulose). If the aim is to recover proteins and polyphenols, these byproducts can be sent to the extraction phase after grinding, used to homogenize particle size.

A byproduct from cereals is lignocellulosic biomass that can be implemented in a biorefinery concept for the production of sugars and further conversion into biofuels or other chemicals. When this is intended, a pretreatment step is required in order to make the enzymes or enzyme-producing
microorganisms accessible to the plant matrix. Many lignocellulosic biomass pretreatment techniques have been developed since early 2000 which involve mechanical pretreatments, such as physical pretreatment (milling), physicochemical pretreatments, and chemical and biochemical pretreatments. For further information about lignocellulosic pretreatment, the reader is referred to the work of Agbor et al. (2011), Sun et al. (2016), and K. Zhang et al. (2016).

The goal of pretreatment is to increase the accessibility of enzymes, to hydrolyze cellulose and hemicellulose into reducing sugars, available in the solid biomass. Chemical and biochemical pretreatments are often the extraction methods for other compounds such as proteins and polyphenols which are discussed in the following section.

**Valuable Compounds Extraction from Agri-food Byproducts**

Proteins, polyphenols, and soluble dietary fiber (oligosaccharides) can be co-extracted from different plant-based side streams. Extraction techniques can be classified into chemical, physical-chemical, and biochemical extraction (Contreras et al., 2019).

Among chemical extraction, solvent extraction and aqueous extraction have been widely applied. Capellini et al. (2017) developed a new method to extract oil from rice bran using safe solvents (ethanol and isopropanol) and mixtures of these solvents with water. The authors obtained around 80% yield of oil with pure ethanol and isopropanol. Additionally, the authors identified that the yield of co-extracted proteins varied up to 20% depending on the solvent, water content, and extraction temperature. The remained protein fraction stays in the defatted rice bran meal; however, the type of solvent and extraction conditions might affect the solubility and functional properties of the proteins.

Traditionally, pectin is extracted mainly from orange peel and apple pomace, in which pectin content ranges from 20 to 40g/100g on dry weight basis (Kulkarni & Vijayanand, 2010), through a process called conventional acid extraction. It makes use of hot water acidified with a mineral or organic acid (e.g., H2SO4); however, the use of strong acid leads to the generation of toxic waste, which should be neutralized before disposal. Additionally, the high temperature and long extraction time can lead to degradation of the pectin structure and decrease in functionality (Rezzadori et al., 2012; Saberian et al., 2017). Other innovative extractive techniques have been studied to recover pectin, such as ultrasound-assisted heating extraction, (Wang et al., 2015), ohmic heating (Saberian et al., 2017), ultra-high pressure (Guo et al., 2012), microwave-assisted extraction (Bagherian et al., 2011), or the use of electric fields (De Oliveira et al., 2015).

Lycopene, which is an important carotenoid with antioxidant properties, has been extracted from tomato pomace and tomato peel using solvent extraction with hexane, ethyl acetate, and ethanol with a yield ranging between 5 (using ethanol) and 120mg/100g (using ethyl acetate) (Calvo et al., 2007). However, due to all the environmental implications of using organic solvents, other technologies have been suggested and successfully applied for the extraction of lycopene. Among these technologies, supercritical CO2 extraction is suggested as a sustainable alternative, which leads to comparable extraction yields than the ones obtained with conventional solvent extraction. Topal et al. (2006) and Kehili et al. (2017) evaluated different operating conditions.
temperature, and pressure, using supercritical CO₂ extraction on tomato peel. The authors obtained 94 and 57% yield of total carotenoids, respectively, using this emerging technique.

Essential oils from citrus fruits have been extracted from the citrus peel by several physical-chemical, mechanical, and thermal techniques, such as cold pressing, solvent extraction, steam distillation, and microwave-assisted extraction (Mahato et al., 2019). Ferhat et al. (2007) extracted essential oils from lemon peels using cold pressing, hydrodistillation, and microwave-accelerated distillation. In hydrodistillation, the plant material is packed and water is added (sufficient amount) and brought to boil. Steam is then introduced and contacted with the plant material and the water; this generates the release of the plant components by hydrodiffusion and hydrolysis. Microwave-accelerated distillation consists on placing the plant material in a microwave reactor and heat it. The internal water in the plant material is then released carrying the essential oils which are then condensed in a condenser outside the reactor (Ferhat et al., 2007). The findings from Ferhat et al. (2007) indicate that microwave-accelerated distillation (MAD) shows comparable yields with hydrodistillation, with a much shorter extraction time (6 times lower). In addition, better antimicrobial activities for oils extracted by MAD were assessed against yeast and gram-negative bacteria, and it is considered and environmentally friendly technique.

Another extraction technique involves the degradation of the cell wall using enzymes. Stoll et al. (2003) developed a process using enzymatic hydrolysis of carrot pomace to recover carotene-rich hydrolysate that can be used as a functional ingredient in, e.g., model beverages (based on apple juice). The process consisted of the application of an enzyme mixture of cellulase and pectinase (cellulolytic and pectolytic activities) to degrade the cell wall of the carrot pomace after milling of this residue. Optimization of the hydrolysis step was the main aspect of the study by Stoll et al. (2003), which was successfully implemented at pilot scale (10 L). The conditions applied in the pilot run were pH 4.0, 50°C, enzyme combination corresponding to 750 ppm of Pectinex Ultra SP-L (Novo Nordisk Ferment), and 750 ppm of Cytolase CL (DSM Food Specialties). The obtained hydrolysate corresponds to 64 mg total carotene per kilogram of hydrolysate. Another well-known application of enzymes is for the hydrolysis of lignocellulosic residues (e.g., sugarcane bagasse) into sugars and later conversion into second-generation ethanol. For this enzymatic reaction, a pretreatment step is required to facilitate the enzyme accessibility to cellulose and hemicellulose presented in the lignocellulosic residues (Mussatto et al., 2010).

Many studies have evaluated protein extraction from agri-food residues. Contreras et al. (2019) provide an excellent review on protein extraction from different agri-food residues, reviewing solvent and aqueous extraction (at alkaline, neutral, and acid conditions), biochemical extraction using enzymes, and physical and physical-chemical methods under dry and non-dry conditions. Regarding protein extraction, it seems that alkaline extraction provides higher yields (up to 95%); however, this method is not selective, so other components such as polyphenols, hemicellulose, and lignin can be co-extracted. Sari et al. (2015) evaluate the protein extraction yield, at alkaline conditions, of different agri-food residues (rape-seed meal, sunflower meal, soybean meal, soybean hull, malt byproducts, among others). The authors identified that biomass composition affects the extraction yield, finding that cellulose and oil can significantly affect the extraction performance. When chemical extraction is low, the use of enzyme-assisted processes might improve protein extraction. For instance, the use of carbohydrate degrading enzymes (cellulase, hemicellulase, xylanase, arabanase, and glucanase) might benefit from the release of proteins from the plant-based matrixes, as they can degrade the cellulose and hemicellulose from the cell wall (Contreras et al., 2019). The use of enzymes can be combined with neutral conditions, preventing the formation of protein-polyphenols complexes and also maintaining the functional properties of the proteins (Fetzer et al., 2018).

As previously mentioned, many valuable compounds are co-extracted and, depending on the application, can be further purified. The following section will outline purification technologies for further separation of valuable compounds mainly in aqueous plant-based extracts.

**Purification of Valuable Compounds from Plant-Based Extracts**

Separation and purification of the valuable components from a plant-based extract can be done using different (conventional and emerging) purification techniques such as adsorption, precipitation or crystallization, membrane separation, aqueous two-phase system separation (ATPS) (Galanakis, 2012), or combinations of these technologies. Adsorption is a promising technology able to purify complex mixtures (plant-based extracts) and also can be operated at mild conditions. Mild operation is often desired in food processing as thermal technologies can generate undesired flavors and change the organoleptic properties of the final products. In addition, adsorption can be a selective process when appropriate adsorbents (polymeric resins, zeolites, activated carbon) are used for capturing the target molecules (products or impurities).

**Protein Purification**

Proteins from vegetable sources are composed of storage proteins (globulin and albumins), structural proteins (ribosomal, membrane proteins), and biological active proteins (lectins) (González-Pérez & Arellano, 2009). Pulses, oilseeds, and vegetables sources are rich in proteins (Fig. 2). Particularly,
Oilseed proteins have been successfully purified by different techniques such as isoelectric precipitation (Akbari & Wu, 2015; Ghodsvali et al., 2005; Karaca et al., 2011; Thiel et al., 2015; Xu & Diosady, 2002) and adsorption (Marshall, 1990; Sewekow et al., 2008; Chung et al., 2005; Bérot et al., 2005; Zhang & Glatz, 1999), being the former one the most applied (González-Pérez & Arellano, 2009). Both techniques are often combined with membrane separation to increase purity. Isoelectric precipitation has been additionally applied in the manufacturing of pulse proteins, after milling of the sources to produce protein-rich flour (Boye et al., 2010). Ultrafiltration can be used to separate proteins (large molecules) from nutrients (small molecules, e.g., polyphenols, sugars, lipids) and antinutritional compounds (phytic acid).

Akbari and Wu (2015) purified napin and cruciferin (storage proteins) from rapeseed meal extract by isoelectric precipitation of cruciferin and subsequent napin purification by removing phytic acid, glucosinolates, and phenolic acids using ultrafiltration (10kDa) and diafiltration, obtaining a total yield of 52% while successfully removing higher than 80% of phytic and phenolic acids. Similarly, Xu and Diosady (2002) and Ghodsvali et al. (2005) successfully purified canola proteins using ultrafiltration and diafiltration before and after isoelectric precipitation of cruciferin fraction. Even though precipitation is widely applied, it has some disadvantages as it can affect the functional properties of the proteins such as solubility and additionally promotes aggregation (González-Pérez & Arellano, 2009). Therefore, the use of mild conditions and technologies such as adsorption could benefit the functionality of the proteins.

Oilseed proteins have been successfully purified by adsorption, decreasing denaturation and obtaining higher purities. Chung et al. (2005) purified the globulin fraction from flaxseed meal using the weak anion exchanger DEAE (diethylaminoethyl) Sephacel, at basic pH (pH 8.6) and mild salt conditions recovering 63% of the total protein content in the meal with high purity. Interestingly, the authors only focused on the recovery of the globulin fraction while an additional minor peak is observed in the chromatogram. This flow through peak corresponded to around 30% of the protein content, which might represent the albumin fraction. Bérot et al. (2005) proposed to purify both protein fractions (albumins and globulins) from rapeseed meal extract using a combination of adsorptive steps. The author used the cation exchange resin, Source 30S, to capture napin and lipid proteins while cruciferin flow through. Cruciferin is subsequently polished by size exclusion (Sephacryl S-300) and napin is polished by HIC (hydrophobic interaction chromatography, Phenyl Sepharose 6 Fast Flow). After polishing both proteins, polyphenols and other antinutritional compounds were removed by dialysis. Purities greater >99% were obtained for both proteins.

Polyphenol Purification

Polyphenol has been purified from the plant-based extract or fermentation broth by means of membrane separation, liquid-liquid extraction (Silva et al., 2018b), preferential crystallization (Silva et al., 2018c), and adsorption (Moreno-González et al., 2020; Sevillano et al., 2014; Silva et al., 2018a; Soto et al., 2011)

Silva et al. (2018b) evaluated the use of liquid-liquid extraction to purify polyphenols from fermentation broth. As one of the critical factors to design a liquid-liquid extraction process is solvent selection, the authors suggested the use of the NRTL-SAC (Nonrandom Two-Liquid Segment Activity Coefficient) thermodynamic model. This model considers the excess entropy and enthalpy to determine the activity coefficient. The NRTL-SAC characterizes any molecule by using four molecular descriptors. The molecular descriptors represent the molecular surface characteristics of the solvent and solute molecules. These descriptors denote hydrophobicity (X), polarity (Y and Y'), and hydrophilicity (Z) (Chen & Song, 2004). The authors, Silva et al. (2018b), use the NRTL-SAC model to predict partitioning into different solvents. The authors suggested different process configurations to purify polyphenols with similar characteristics. In another study, Silva et al. (2018c) used preferential crystallization (commonly used for enantiomer purification) to separate naringenin and trans-resveratrol (chemically related polyphenols). This study showed an alternative technique besides adsorption for recovering similar compounds; however, the solution needs to be concentrated and cooled to induce supersaturation and later crystallization.

Adsorption has been widely applied for recovery of phenolic compounds from liquid food streams. The studies performed by Pickardt et al. (2015) and Weisz et al. (2010) evaluated the adsorption of phenolics onto a polymeric resin and an ion exchange resin (XAD16 and Lewatit S 6328) using sunflower meal extract. Both studies aimed to recover proteins; however, they suggested the co-recovery of polyphenols compounds as a possibility to improve economic potential of the process. Thiel et al. (2015) evaluated the recovery of proteins, sinapic acid, and phytic acid from rapeseed meal extract. After extraction, purification was assessed with two processes using a β-zeolite in the H+ form (SiO2/Al2O3 ratio of 150:1) and the anion exchanger Purolite A200. Both purification processes accomplished the separation of the three components of interest. Ferri et al. (2011) studied the recovery of hydroxycinnamic acids from olive mill wastewater, using a synthetic mixture of ten polyphenols, which are the most representative ones of this side stream. The authors of this work also used macroporous food grade resins and a weak anion acidic resin.
exchanger (XAD4, XAD7, XAD16, IRA96, and ENV+). An overall of 90% polyphenol recovery was found when desorbing with ethanol. Similarly, Moreno-González et al. (2020) evaluated the recovery of sinapic acid from rapeseed meal extract using a model feed system. Adsorption of sinapic acid on the polymeric resin FPX66 resulted to be selective. They demonstrated that other components such as sugars, glucosinolates, and phytic acid (often presented in food matrices) poorly interact with the resin which benefits sinapic acid capture. At last, Schieber et al. (2003) developed a process using apple pomace extract to recover polyphenols and pectin. Phenolics are purified by adsorption on XAD16 and subsequently eluted using methanol. Solvent was evaporated and polyphenols were freeze dried.

**Polysaccharides Purification**

Pectin purification from citrus and fruit extracts (peels and pomace) can be accomplished by solvent precipitation, ion exchange chromatography, and dialysis (Lampitt et al., 1947), among others technologies. Besides bioactive phenolics from apple pomace, Schieber et al. (2003) purified pectin by alcohol precipitation of the pectin-containing effluent from the adsorption column where phenolics were captured.

Solvent precipitation with ethanol was applied by Galanakis et al. (2010) to purify dietary fibers from olive mill wastewater. After concentration of the wastewater (3.5-fold) and subsequent liquid-liquid extraction using ethanol/acid and ethanol/water mixtures, dietary fibers were precipitated by an increase of ethanol concentration up to 85mL/100 mL. Precipitated fibers were deflated with acetone and dried. As mentioned earlier, olive mill wastewater also contains significant amount of polyphenols. The suggested purification process by the authors allows the co-recovery of dietary fibers and polyphenols. As the fibers precipitate in ethanol, phenolics are soluble in this solvent. The recovered dietary fiber was further characterized, finding that the soluble fraction is rich on pectin polysaccharides while the insoluble fraction is rich of glucose, xylose, galacturonic acid, and rhamnose.

Ion exchange chromatography has been successfully implemented for sucrose purification from beet molasses (Ganetos & Barker, 1992) and separation of fructose and glucose (Azevedo & Rodrigues, 2001; Hashimoto, 1983, 1987). Chilamkurthi et al. (2012) assessed the different cation exchangers in the form of Na⁺, H⁺, K⁺, and Ca²⁺, to evaluate the separation of arabinose and sugars. The model components used in this study were galactose, glucose, arabinose, lactose, and sugar acid. Their findings indicate that appropriate separation of the different components is accomplished using the cation exchangers in the form of Ca²⁺ and K⁺.

Another purification method for polysaccharides like sugars, glucose, is crystallization. As crystallization occurs when the solution is supersaturated, a pre-concentration is required. The shape and size of crystal will determine the subsequence steps to separate them from their mother liquids. These steps will involve filtration and drying (Berk, 2013a).

**Flavor Ingredients Recovery/Purification**

Natural flavors, which are generally obtained from plant or animal sources, are widely used in the food sector mainly in beverages. Saffarianpour and Ottens (2018) provide an excellent review on the different technologies applied for purifying these types of products, including distillation, pervaporation, and adsorption.

Additionally, adsorption has been applied for capturing of impurities (e.g., off-flavors) instead of the valuable products. Gernat et al. (2020) successfully decreased the concentration of wort off-flavors (2-methylbutanal, 3-methylbutanal, methional, 2-methylpropanal, and furfural) in alcohol-free beer using zeolites at pilot scale (150 L). The authors evaluated other adsorbents such as polymeric resins; however, it was proved that resins are able to adsorb other flavor components, affecting the quality of the final product, as the aim was only to remove wort flavors. Off-flavor components are mainly composed of hydrophobic organic molecules (aldehydes, ketones, carboxylic acids). The results from Gernat et al. (2020) introduced the possibility of removing these components from food liquid streams by using zeolites to increase selectivity to aldehydes or by using hydrophobic resins for overall off-flavors removal.

**Final Product Formation**

The last step toward the manufacturing of a product from a side stream of a food process corresponds to formulation, defined here as the ultimate product form (emulsions, powders, pieces, liquid, among others).

**Food Powders/Solid Extracts**

Drying technology is mostly used for food preservation, as minimization of moisture content in food products inhibits microbial growth. However, drying can also be used to improve the physical properties of a product (Smith, 2011), for instance, spray drying is used to dry liquid food and form a powder product. Among the different types for dryers utilize by food industry, pneumatic drying, spray drying, drum drying, tray drying, and freeze drying are applied in formulation.

Spray drying has been applied to concentrate food liquids such as coffee, milk, and juices. This technique is also used for encapsulation of food ingredient, such as flavors and whole food substances (e.g., chocolate), where materials can be encapsulated as single particle structure (ingredient surrounded by a matrix wall) or in an aggregate structure (ingredient particles fixed in a matrix) (Desai & Jin Park, 2005). One of the main
advantages of spray drying is that it produces a stable particulate solid product from a liquid in a one continuous step. However, wall matrix materials are limited, being the most applied maltodextrins, hydrophobically modified starch, gum acacia (Desai & Jin Park, 2005), whey, gelatin and sucrose (Anandharamakrishnan & Ishwarya, 2015). Spray drying seems to be a suitable technique in the final formulation of a product from a side stream of food (plant extract, wastewater) as these streams are mainly liquid streams. In addition, spray drying is relatively economical and suitable for heat-sensitive materials (Anandharamakrishnan & Ishwarya, 2015).

Freeze drying is mostly applied in the pharmaceutical field and also in the production of expensive biological products when product quality is the most important attribute of the product. Its application in the food sector for the recovery of products might be limited to high-value products such as proteins, enzymes, and heat-sensitive products as flavors. The principle consists on freezing the feed on a chamber where vacuum is applied. Dehydration occurs by sublimation, mostly of water which is recovered by mechanical vacuum pumps (Barbosa-Cánovas et al., 2005). Note that freeze drying is usually operated in batch mode, as it requires a drying chamber where the product is frozen and longer drying times than other techniques such as spray and drum drying, which limits its application to small throughputs. The product form obtained from freeze drying is pieces, which are usually grounded after processing (Barbosa-Cánovas et al., 2005).

Desobry et al. (1997) evaluated the drying process of β-carotene encapsulation using freeze drying, spray drying, and drum drying. The authors demonstrated that the characteristics of the dried β-carotene obtained from drum drying possess similar characteristics than the one obtained with the other two methods. In addition, during storage, the authors identified that drum drying gave higher product retention comparing to the other two methods. The solid obtained in the shape of sheets can be grounded to different particle sizes.

Other types of dryers that can be used for product formulations are tray dryers, bin dryers, rotatory dryers, and pneumatic dryers. Particularly these dryers are used (but not limited) when the feeds stream is composed of solids (particulate, grains, granulated materials, precipitates, or crystals). Bin and tray dryers are operated in batch mode while rotatory and pneumatic dryers can be operated in continuous mode (Berk, 2013b).

Selection of drying technique will highly depend on the desired product characteristics and the processing volume.

Food Byproducts Processing Examples

As previously mentioned, the recovery and purification of valuable compounds from food side streams (bran, wastewater, oilseed meal, fruit pomace, among others) is dictated by the products to be recovered and the nature of the source material. In the following paragraphs, examples of purification processes are described based on the previously discussed techniques and literature.

Galanakis et al. (2010) co-recovered dietary fiber from olive mill wastewater (Fig. 4a). The suggested process consisted in a pre-concentration step of the wastewater, which might help to increase the extraction yield. Valuable compounds were extracted from the wastewater using ethanol by adding 5 mL of ethanol and 1 g of citric acid and completed 100 mL of mixture with the concentrated wastewater. Extraction was done at 80°C for 10 minutes. Dietary fiber was precipitated by contacting the extraction mixture in 95% (v/v) ethanol up to a concentration of 85% (v/v) and boiled for 10 minutes. Precipitated dietary fiber was filtrated and washed with acetone and chloroform to remove remained oil and dried for further analysis. The discarded liquid from the precipitation is rich in polyphenols. The authors found a concentration of 1.25g/L which is similar to the concentration of phenolics found in extra virgin oil. The phenolic rich fraction might contain different types of phenolic compounds, mainly hydroxycinnamic acids. These phenolic compounds could be further separated from the liquid by crystallization and dried as suggested by Silva et al. (2018b). Recovery of solvents could be accomplished by distillation and can be recycled to the extraction system which might additionally benefit process economics.

When using oilseed meals as byproducts, proteins, dietary fibers, and polyphenols could be purified. Aqueous extraction assisted with salt could be applied as an alternative method, instead of solvent extraction or alkaline extraction. Separation of small molecules (sugars and polyphenols) from the large molecules (proteins) can be accomplished using membranes (Ghodsvali et al., 2005; Xu & Diosady, 2002). Isolate of both proteins could be further processed by further membrane concentration and freeze drying; however, each protein has specific applications. Napin is a basic protein stable at a wide pH range and holds foaming properties while cruciferin possesses emulsifying, gelling, and binding properties (Aider & Barbana, 2011; Wanasundara et al., 2016). Purification of both proteins by ion exchange chromatography has been suggested (Moreno-González et al. (submitted manuscript)), where napin is bound to the resin while cruciferin flow through. The purified protein fractions after chromatography can be sent to another membrane unit to remove buffer salts and dried. The permeate, from the small molecules’ separation, contains the remained co-extracted components, polyphenols, sugars, phytic acids, and glucosinolates. Moreno-González et al. (2020) suggested capturing the polyphenols compounds using a food grade macroporous resin (FPX66). The authors demonstrated that the other components, sugar, phytic acid, and glucosinolates poorly interact with the resin. The obtained sinapic acid (mayor phenolic in rapeseed meal)
fraction, recovered using ethanol/water mixture (Ferri et al., 2011; Moreno-González et al., 2020; Silva et al., 2018a), could be crystallized (Silva et al., 2018b), washed, and dried (Fig. 4b). As with the previous example, solvent could be additionally recovered by distillation. The flow through fraction from the adsorption column contains sugars, phytic acid, and glucosinolates. Phytic acid is well known as an antinutrient compound due to its ability to bind to mineral ions such as magnesium and calcium, therefore is often removed from food products. However, sugars are valuable compounds and further purification of sugars could be accomplished by chemical precipitation of phytic acid. The remained solid fraction from the aqueous extraction is rich in dietary fibers hemicellulose, cellulose, and lignin, which could additionally be processed to obtain reducing sugars for biofuels.

Following some of the strategies suggested previously, processing of fruit/tuber pomace (Fig. 5a) could start with the acid extraction of the polyphenols and soluble carbohydrates, which was applied for apple pomace by Schieber et al. (2003) and could be extended to other type of pomace such as carrot or tomato. The extract contains polyphenol components and soluble sugars which could be purified by adsorption using food grade hydrophobic polymeric resins such as XAD16 and FPX66 as soluble sugars might poorly interact with the hydrophobic matrixes (Ferri et al., 2011; Moreno-González et al., 2020; Schieber et al., 2003). After polyphenol elution, Schieber et al. (2003) recommended to evaporate the solvent, which could be recovered by distillation and remove the remained water by freeze drying. Among the polyphenols that could be potentially recovered, there is lycopene from tomato pomace (Lu et al., 2019), carotene from carrot pomace (Stoll et al., 2003), and phloridzin and chlorogenic acid from apple pomace (Schieber et al., 2003). The flow through obtained liquid from the adsorption
is rich in pectin, which can be purified by alcohol precipitation, filtrated or centrifuged, to remove liquid, and finally dried.

Lignocellulosic biomasses from food byproducts such as cereal bran could be processed to recover antioxidants (polyphenols), proteins, and fermentable sugars (Fig. 5b). As previously mentioned, cereal brans are rich on phenolics and dietary fibers. Aqueous extraction (alkaline or acid) of proteins and polyphenols is then performed. Note that alkaline conditions could oxidize the phenolic compounds or form protein-polyphenols complexes; milder conditions, like neutral extraction, are therefore preferred. Solid-liquid separation between the extract and the insoluble fiber can be done by centrifugation of filtration. Purification of the plant-based extract can be done by isoelectric protein precipitation as recommended by Celiktas et al. (2014), and polyphenols could be further purified by adsorption (hydrophobic polymeric resins) (Ferri et al., 2011; Moreno-González et al., 2020; Schieber et al., 2003; Silva et al., 2018a). The protein precipitate could be filtrated and dried while phenolics could be crystallized and dried. The solid fraction from the aqueous extraction could be further treated by liquid hot water pretreatment for autohydrolysis of hemicellulose followed by the hydrolysis of the plant fibers using enzymes to produce fermentable sugars that can be later converted into biofuels or other biochemicals.

**Economic Considerations**

This work has extensively discussed the technical feasibility of recovering valuable compounds from food side streams.
However, for large-scale implementation, economic feasibility is as important as technological potential.

Economic considerations include three main factors: (1) estimation of capital investment, (2) operating cost, and (3) profitability analysis. These factors can be assessed using well-known methodologies described elsewhere (Harrison et al., 2015; Seider et al., 2010; Towler & Sinnott, 2013). Regarding capital investment, one of the main considerations is the cost of the land when new facilities are built. Therefore, careful considerations, such as proximity to the facilities where the side stream is generated, should be taken into account. Proximity or even installing new equipment in the facilities will consequently reduce transportation cost and therefore increase economic potential. Moreover, the aim is to have low CAPEX (capital expenditure) and OPEX (operating cost).

Knowing that voluminous streams are typically produced in the food sectors, recovery and purification of the valuable compounds seems to be an attractive option that can substitute the cost associated to their disposal which would potentially be economically favorable, as, when treated in the same facilities, might reduce raw material costs (operating cost). In addition, minimization of additional waste streams should also be considered by recovering several multiple products from a single stream. Furthermore, this might benefit economic feasibility as more products could be sold, reducing or even removing the cost associated to them when treated as waste.

Moreover, implementation of continuous operation (often desired by the food sector) and counter current operation might lead to more efficient operation and cost reductions. In terms of CAPEX, this might lead to smaller equipment, while for OPEX, considering as an example, continuous chromatography, this might reduce buffer consumption and adsorbent cost (Steinebach et al., 2016). Semicontinuous chromatography such as simulated moving bed (SMB) has shown great potential in different industry fields, such as petrochemical, pharmaceutical, biopharmaceutical, and food (Ganetsos & Barker, 1993).

At last, it is important to consider that the selling cost of the products (profitability analysis) should be competitive with the market and process considerations such as process intensification and heat integration should be carefully assessed during process design.

Techno-economic evaluation (TEE) combined with life cycle assessment (LCA) has been suggested to assess biorefineries (Brown et al., 2014). This combination is also applicable to assessing the processing of food side streams to recover valuable products. LCA estimates the environmental impact associated to process operation, from raw materials to disposal and/or recycle, while TEE assesses technical feasibility and economic potential. By combining both assessments, the findings of each discipline are enhanced (Brown et al., 2014) which might help on decision-making to mitigate harmful environmental implications of processing food side streams.

Future Prospects

Multiple valuable products can be recovered from a single food side stream depending on the selected processing steps and the source origin (strong, soft, and wastewater). Technology Readiness Level (TRL) Index, introduced by NASA (Héder, 2017), can be used to assess the stage toward large-scale implementation where certain technology is located. The TRL index scale goes from 1 to 9, where TRL 1 concerns to fundamental concept/research, TRL 2–4 to technological research, TRL 5–8 to product demonstration, and TRL 9 to implementation (Roque et al., 2020).

For processing lignocellulosic biomass (strong structures), selection of a suitable extraction technique is crucial in order to recover several products. Different extraction techniques can affect the properties of the products to be recovered. Even though alkaline extraction is the most reported method, it also could promote protein denaturation and phenolic acid oxidation, which consequently reduce product yield and quality. Even though extraction is a well-established technology (TRL 9), the use of milder conditions such as neutral extraction, other extraction systems such as aqueous two-phase systems (ATPS), or alternative techniques such as supercritical CO2 (TRL 5-8) should be further studied and optimized for this step. However, this implicates higher R&D associated cost and these technologies are not yet implemented in large scale.

Among other separation techniques, adsorption is an attractive alternative for capturing of valuable products such as proteins and polyphenols due to its simplicity, high efficiency, ease to scale up, and selectivity when appropriate adsorbents are available. Particularly for polyphenols adsorption, it has been demonstrated that food grade polymeric resins (e.g., XAD16, FPX66 from the Amberlite series) are highly effective. Often elution of the bound compounds concentrates them which facilitate their further processing by crystallization/precipitation and drying. Adsorption could also be used for removal of unwanted compounds in a liquid stream such as off-flavors (Gernat et al., 2020) and colorants, as already applied for de-bittering of fruit juices. Even though packed bed chromatography is a well-established technology (TRL 9), better understanding of the interactions between the adsorbent and the bound components might accelerate the implementation of more adsorptive processes in the food sector. Moreover, and as previously mentioned, continuous operation (TRL 5–8), not only with SMB systems (simulated moving bed systems) but with emerging systems such as periodic countercurrent chromatography (PCC), might benefit techno-economic potential.
Fruits and vegetables residues (soft structures) are low in protein content but rich in polyphenols and carbohydrates such as pectin. As with the residues from cereal, oilseed, and pulses, extraction methods will greatly influence the following purification steps. Alcohol extraction is a very effective technique for polyphenols as it utilizes the higher solubility of these types of components in alcohols. Even though solvents could be recovered by distillation, the environmental implications that they represent encourage the need of utilized alternative extraction techniques. Great progress has been accomplished with supercritical fluid extraction, which additionally provides selectivity toward the extracted compounds (Saffarionpour & Ottens, 2018). However, large-scale implementation has not been fully established and more effort is needed in order to guarantee safety during operation.

The last stage of the processing will be related to the ultimate product form, which commonly is a solid form. Therefore, water of other solvents should be additionally removed by drying. Spray drying, drum drying, and freeze drying are the few drying technologies able to dry liquid streams. This work describes a structure approach of the different process steps required to recover multiple value-added compounds from a single agri-food streams. This approach combined with TEE, LCA, and TRL might facilitate process design of new food side streams. Recovered components with high purity level can be applied in pharmaceutical and cosmetic sectors. For the food sector, where functionality (emulsification, gelling formation, thickening, viscosity) is given by combining several ingredients, high degree of purity might not be needed. Limited studies are available (Geerts et al., 2017; Karefyllakis et al., 2019; Kornet et al., 2020), exploring this new way of thinking, and they have shown that the properties of the final products are not significantly changed.

Concluding Remarks

This review presents a structure approach of the different process steps required to recover valuable components from food side streams. It demonstrates that a multitude of valuable compounds can be recovered from a single side stream including, in most of the cases, polyphenols, carbohydrates (dietary fibers), and proteins. The recovery of multiple products depends on the source origin (strong and soft matrices and wastewater).

It is clear that to fully isolate the extracted compounds, several separation technologies will be combined and a detailed life cycle assessment (LCA) and techno-economic evaluation (TEE) should be performed to evaluate feasibility and profitability.

As previously mentioned, the application of the different technologies will depend on their level of technological readiness to large-scale implementation (TRL) and its suitability to continuous operation. Moreover, the processing of plant-based side streams should be cost-effective, and keeping the functionality of the products (e.g., nutritional value of proteins and antioxidant activity in polyphenols) in order to be successfully applied in pharmaceutical, cosmetic, and food products. The presented approach might improve the valorization of industrial food side streams which could be beneficial in terms of sustainability.

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