Abstract

Plant molecular farming depends on a diversity of plant systems for production of useful recombinant proteins. These proteins include protein biopolymers, industrial proteins and enzymes, and therapeutic proteins. Plant production systems include microalgae, cells, hairy roots, moss, and whole plants with both stable and transient expression. Production processes involve a narrowing diversity of bioreactors for cell, hairy root, microalgae, and moss cultivation. For whole plants, both field and automated greenhouse cultivation methods are used with...
products expressed and produced either in leaves or seeds. Many successful expression systems now exist for a variety of different products with a list of increasingly successful commercialized products. This chapter provides an overview and examples of the current state of plant-based production systems for different types of recombinant proteins.

**Keywords**
Molecular farming · Recombinant protein expression · Therapeutic proteins · Transient expression · Bioreactor

## 1 Introduction

Photosynthesis or use of simple growth media provides clear advantages for using plants to produce biomass and protein products at low cost [1–3]. The majority of posttranslational modifications important for many complex eukaryotic proteins can be performed by plants, whose species diversity offers variety in production platforms from in vitro cultures through field crops, all within established regulatory guidelines. Of particular benefit is that plants cannot harbor human and animal pathogens that can plague in vitro mammalian production systems, so plants provide major advantages in product safety, especially for therapeutics [4, 5]. Such safety advantages also reduce purification costs and minimize production shutdowns and facility decontamination, thereby affecting patient/customer demand. Costs to purify plant-produced proteins are comparable to microbial or mammalian cell culture systems, but plants offer key advantages with lower up-front capital costs and potential economies of scale. Direct use of cultivated plant cells, whole or minimally processed plants, or plant parts (e.g., seeds, dried leaves) is in development for industrial/bioenergy applications as well as for therapeutics and vaccines, thereby further reducing costs of recombinant proteins [6–12]. Boosting protein yields is always a challenge for economic feasibility [2, 5] as are regulatory considerations, some of which are unique to plant bioproduction systems [1, 13].

Plant expression platform diversity includes: whole plants, suspension cells, hairy roots, moss, duckweed, and microalgae. There are strengths and weaknesses to each platform with selection often determined by type of recombinant protein, market, scale, cost, and up and downstream processing constraints of the specific protein product. Within each platform is also a diversity of plant species that can host the protein product. Platform diversity provides flexibility in expressing novel recombinant proteins, enables customizing and meeting scale-up needs, and provides opportunity for oral-based delivery of proteins. However, such diversity also may complicate regulatory compliance, which prefers more uniformity. Here we summarize prospects and challenges associated with each type of plant production platform and production strategy, and where appropriate, comments are provided on important regulatory issues and progress toward commercialization.
2 Types of Recombinant Proteins Produced by Plants

Plant-made recombinant proteins are generally categorized into three classes, usually based on function and/or application: biopolymers, industrial proteins/enzymes, and therapeutic proteins. Examples of each of these groups are briefly described in the following sections.

2.1 Biopolymers

Although plants are expert at bulk production of biopolymers such as cellulose and starch, here the focus is on recombinant protein-based biopolymers such as elastin-like polypeptides (ELPs), spider silk proteins, collagens, and plant gums (see review [14]). The ELPs that are repetitive pentapeptide sequences (VGVPQ) mainly serve as thermally responsive tags for non-chromatographic purification of recombinant proteins [15]. ELP tags significantly enhanced production yield of different recombinant proteins in plant leaves [16, 17]. Collagen and spider silk proteins (spidroins) serve as two interesting examples.

Collagen is a critical extracellular matrix (ECM) material and connective tissue in animals; commercial supplies are typically extracted from cadavers or animal sources, so a safer source is desirable [18]. Large-scale production of recombinant human collagen type I (rhCOL1) in tobacco is now providing low-cost functional biocompatible ECM that is safer than cadaver or animal sources for use in creating scaffolds for tissue engineering, skin, and wound healing [19, 20].

Spidroins are modular fibrous proteins with highly repetitive amino acid sequences consisting mainly of glycine and alanine [21, 22]. Silk fibers spun from these spidroins have exceptional flexibility, elasticity, and toughness – three times as strong as Kevlar and five times as strong as steel [23]. Plant production is more efficient and cheaper than microbial recombinant spidroins. Transgenic tobacco and potato plants expressing Nephila clavipes synthetic genes of dragline spidroin have yielded recombinant silk proteins up to 2% of total soluble protein (TSP) in the ER [22]. Arabidopsis dragline spidroins reached 18% of TSP in seeds [24]. Challenges remain in developing manufacturing technology for spinning fibers into more useful products [25].

2.2 Industrial Proteins/Enzymes

Industrial proteins are defined as commodity chemicals used in very large quantities and thus must be produced at very low cost [26]. Transgenic field plants are particularly attractive for producing industrial proteins/enzymes because agricultural production is low cost, protein products stored in specific organs such as seeds are stable, scale-up is easy and relatively fast, and crude plant materials can often be used directly in industrial processes [6, 7]. Industrial proteins of most interest are hydrolases, including glycosidases (e.g., cellulase, α-amylase, and β-glucuronidase...
(GUS)) and proteases (e.g., trypsin). The first company to develop and commercialize plant-based recombinant proteins/enzymes, with GUS and avidin being their first two commercialized products, was ProdiGene Inc. (formerly in College Station, TX), which is unfortunately now out of business [3, 7, 27].

Corn seed is considered an ideal platform for industrial protein/enzyme production because this plant has the largest annual grain yield and relatively high seed protein content (10%), offering the highest potential recombinant protein yield per hectare [28]. Regulation is a major hindrance to using plant-made industrial proteins/enzymes because a large acreage of transgenic plants is needed. Use of a food/feed crop for nonfood/feed products may also meet political resistance when, as projected, global food supplies become limited, therefore requiring new solutions.

### 2.3 Therapeutic Proteins

Biopharmaceutical sales are projected to reach $US 278.2 billion by 2020. Although transgenic plants can produce fully functional mammalian proteins, including blood proteins, vaccine antigens, monoclonal antibodies (mAbs), cytokines, therapeutic enzymes, growth factors, and growth hormones [5, 29–31], bioactivity requires proper folding, disulfide bond formation, subunit assembly, and often proteolytic cleavage and/or glycosylation. Plants can produce lower-cost, safer therapeutic proteins than mammalian cells [30] and may be the only production system available, e.g., for production of secretory antibodies (sIgAs) [2]. These recombinant therapeutic proteins are produced using many different plant-based platforms including cells and field crops; the most common production species is tobacco. Protein yields have reached well over 25% of TSP [32] 247 mg L⁻¹ [33].

Plants successfully perform N-linked glycosylation of proteins at the signature recognition motif (N-X-S/T) with subsequent processing in the Golgi complex. As a result, plant glycans differ from those found in mammalian cells. Using plants as hosts to produce therapeutic glycoproteins results in addition of plant-specific xylose and α-1,3-fucose sugars; these may alter bioactivity or immunogenicity in humans [34, 35]. The β-1,4-galactose or sialic acid residues synthesized in mammals are not naturally produced in plants [31, 35–37], so RNAi strategies have been used to engineer more humanlike glycosylation machinery in plants by knocking down fucosyl- and xylosyltransferases in plant transgenic lines expressing a human or chimeric β-1,4-galactosyltransferase [38–40]. Furthermore, tobacco plants infiltrated with *Agrobacterium tumefaciens* produced human antibodies with humanlike N-glycans when co-expressed with a chimeric human β-1,4-galactosyltransferase [41]. The first human-injected therapeutic with greatest clinical experience (Protalix’s taliglucerase alfa; discussed further below) did not trigger significant patient antibody production [42–44]. Indeed, plant-specific posttranslational modifications (PTMs) may offer opportunities for producing novel recombinant proteins with enhanced function and efficacy as well as biosimilars [45]. “Second generation” plant-made pharmaceutical proteins are emerging, wherein therapeutic targets are specifically engineered to enhance or produce new therapeutic proteins that integrate
novel motifs or fusion to facilitate protein assembly, delivery, trafficking, protein stability, serum longevity, or protein solubility in either the production host or the target organism (e.g., [31, 46–49]).

3 Platforms for Plant Expression

Advances in plant molecular engineering technologies have expanded the diversity of plant bioproduction platforms, ranging from cell and tissue cultures under sterile and contained conditions to whole plants grown under glass or in the field [2]. These bioproduction platforms can be classified as: (1) in vitro culture systems including cell suspensions, hairy roots, and moss protonema, (2) aquatic plants including duckweed and microalgae, and (3) whole plants using both stable and transient expression. The stable expression of whole plants also includes leaf- and seed-based systems. Characteristics of each platform and their strengths and weaknesses are described here. A comparison of the cost, applicability, time needed for production, scalability, and regulatory compliance of different platforms is shown in Table 1.

3.1 In Vitro Culture Systems

Plant biomass (e.g., suspension cells, hairy roots, and moss) can be propagated in confined bioreactors under sterile conditions for large-scale production of recombinant proteins. In vitro culture allows for precise control over growth and protein production, batch-to-batch product consistency, and a production process aligned with current good manufacturing practices (cGMP) [5]. Due to its relatively high cost of production, in vitro cultures are often used to produce high-value protein therapeutics. Compared with the other plant-based platforms, in vitro cultures are more acceptable to the pharmaceutical industry with fewer regulatory and environmental concerns [50]. Like other bioreactor-based culture systems, scalability of in vitro cultures is limited by bioreactor capacity. However, because recombinant proteins can be secreted into culture media, downstream processing of recovery and purification of the proteins becomes less expensive than from whole plants [51]. Indeed, the first licensed pharmaceutical protein derived from plants for human use – taliglucerase alfa (Elelyso™) – was produced in carrot suspension cells. Examples of representative recombinant proteins produced using in vitro cultures are shown in Table 2.

3.1.1 Plant Cell Suspensions

Undifferentiated clusters of plant callus can be dispersed and propagated in a liquid medium to generate stable cell suspension cultures that retain the same production capacity as whole plants. The production of recombinant proteins in plant cell culture was first demonstrated in 1990 with the expression of a human serum albumin in tobacco cells [78]. Since then, a wide array of biologically active proteins, including antibodies, vaccine antigens, growth hormones and factors,
Table 1 Comparisons of different plant-based bioproduction platforms

| Platforms                          | Viable species                                      | Time for production<sup>a</sup> | Types of proteins applicable          | Scalability   | Production cost<sup>b</sup>       | Regulatory compliance |
|------------------------------------|----------------------------------------------------|---------------------------------|---------------------------------------|---------------|-----------------------------------|-----------------------|
| **In vitro culture system**        |                                                    |                                 |                                       |               |                                   |                       |
| Plant cell suspensions             | Tobacco (Nicotiana tabacum cv. BY-2), carrot, rice | 7–14 days                       | Therapeutic proteins                  | LBC<sup>c</sup>, as high as 100,000 L | Medium              | Easy                  |
| Hairy roots                        | N. tabacum                                          | 14–30 days                      | Therapeutic proteins                  | LBC, as high as 20,000 L             | Medium              | Easy                  |
| Moss                               | Physcomitrella patens                               | 14–30 days                      | Therapeutic proteins                  | LBC, as high as 500 L                | Medium              | Easy                  |
| **Aquatic plants (contained system)** |                                                  |                                 |                                       |               |                                   |                       |
| Duckweed                           | Lemna sp., Spirodela sp.                            | 20–40 days                      | Therapeutic proteins; industrial enzymes | LBC, as high as 10,000 L            | Low                 | Moderate              |
| Microalgae                         | Chlamydomonas reinhardtii, Dunaliella salina       | 14–30 days                      | Therapeutic proteins                  | LBC, as high as 10,000 L            | Low                 | Easy                  |
| **Whole plants (leaf-based and seed-based system)** |                              |                                 |                                       |               |                                   |                       |
| Stable transgenic plants           | Corn, soybean, safflower, rice, tobacco             | 3–6 months                      | Therapeutic proteins; biopolymers; industrial enzymes | Unlimited field culture | Very low              | Difficult             |
| Transient plants                   | N. benthamiana, lettuce                            | 2–7 days                        | Therapeutic proteins                  | Greenhouse limited                  | Low                  | Easy                  |

<sup>a</sup> The time required to accumulate maximum amounts of recombinant proteins in a culture system after planting or bioreactor inoculation

<sup>b</sup> Compared with mammalian cell culture platform

<sup>c</sup> LBC limited by bioreactor capacity
### Table 2  Example of representative recombinant proteins produced via in vitro cultures

| Recombinant proteins | Host plant species | Promoter       | Protein yields  | References |
|----------------------|--------------------|----------------|-----------------|------------|
| **Plant suspension cells** |                    |                |                 |            |
| Anti-HIV antibody 2G12 | *N. tabacum* cv BY-2 | CaMV35S | 12 mg L⁻¹ | [52]        |
| Human antibody M12    | *N. tabacum* cv BY-2 | CaMV35S | 107 mg L⁻¹    | [53]        |
| Anti-hepatitis B surface antigen (HBsAg) mAb | *N. tabacum* cv BY-2 | CaMV35S | 15 mg L⁻¹ | [54]        |
| Human growth hormone (hGH) | *N. tabacum* cv BY-2 | CaMV35S | 35 mg L⁻¹ | [49]        |
| Human interferon α2b (hIFNα2) | *N. tabacum* cv BY-2 | CaMV35S | 28 mg L⁻¹ | [48]        |
| Human α-1-antitrypsin (rAAT) | *O. sativa* | RM3D | 247 mg L⁻¹ | [33]        |
| hGM-CSF               | *O. sativa*        | RM3D          | 129 mg L⁻¹     | [56]        |
| Glucocerebrosidase   | *D. carota*        | CaMV35S       | N/A            | [57]        |
| α-Galactosidase-A (PRX-102) | *D. carota* | CaMV35S | N/A | [58]        |
| Human interleukin-12 (IL-12) | *O. sativa* | RM3D | 31 mg L⁻¹ | [59]        |
| Human serum albumin (hSA) | *O. sativa* | RM3D | 45 mg L⁻¹ | [60]        |
| Enhanced GFP          | *N. tabacum* cv BY-2 | CaMV35S | 125–131 mg L⁻¹ | [61]        |
| **Hairy roots**       |                    |                |                 |            |
| Human antibody M12    | *N. tabacum*       | CaMV35S       | 5.9 mg L⁻¹     | [62]        |
| Human antibody H10    | *N. tabacum*       | CaMV35S       | 2–3 mg L⁻¹     | [63]        |
| Hepatitis B surface antigen (HBsAg) | *Solanum tuberosum* (var. *Kufri bahar*) | (Aocs)AmasPmas | 97.1 ng g⁻¹FW | [64]        |

(continued)
| Recombinant proteins                      | Host plant species | Promoter       | Protein yields<sup>a</sup> | References |
|------------------------------------------|--------------------|----------------|-----------------------------|------------|
| 14D9 murine IgG1                         | *N. tabacum*       | CaMV35S        | 64 mg L<sup>−1</sup>        | [65]       |
| Human epidermal growth factor (hEGF)     | *N. tabacum*       | CaMV35S        | 2 ug g<sup>−1</sup>FW       | [66]       |
| Thaumatin                                | *N. tabacum*       | CaMV35S        | 2.6 mg L<sup>−1</sup>       | [67]       |
| GFP                                      | *Brassica rapa* (turnip) | CaMV35S      | 120 mg L<sup>−1</sup>       | [68]       |
| Human tissue plasminogen activator (tPA) | *Cucumis melo L cv Geumssarageuncheon* or *N. tabacum* | CaMV35S      | 798 ug g<sup>−1</sup>FW or 900 ng mg TSP | [69, 70] |
| Moss                                     | *Physcomitrella patens* | PpUbq1<sup>b</sup> | 250 µg g<sup>−1</sup>DW     | [71, 72]  |
| Vascular endothelial growth factor (VEGF) | *P. patens*        | Ppact5-P<sup>c</sup> | 23.2 µg g<sup>−1</sup>DW (secreted); 25.8 µg g<sup>−1</sup>DW (intracellular) | [75]       |
| Erythropoietin (EPO)                     | *P. patens*        | Pca200P<sup>d</sup> | 3.7 µg g<sup>−1</sup>FW    | [76]       |
| Complement factor H                      | *P. patens*        | N/A            | N/A                         | [77]; www.greenovation.com |
| Multi-epitope fusion protein (HIV vaccine)| *P. patens*        | N/A            | N/A                         | [77]; www.greenovation.com |

<sup>a</sup>Fresh weight, DW dry weight, TSP total soluble protein, NA date not available

<sup>b</sup>Chimeric super-promoter consisting of three copies of the octopine synthase activator (*Aocs*) and one copy of the mannopine synthase activator (*Amas*) located upstream of the mannopine synthase promoter (*Pmas*)

<sup>c</sup>*Ppact5-P*: *P. patens* actin<sup>5</sup> gene promoter

<sup>d</sup>*PpUbq1*: *Physcomitrella* ubiquitin gene-derived promoter

*Pca200P*: *P. patens* carbonic anhydrase 200 promoter
cytokines, and therapeutic enzymes, have been successfully produced in plant cell culture. See recent reviews [5, 50, 79].

Plant suspension cultures integrate the merits of whole plant production with those of microbial fermentation and mammalian cell culture [80, 81]. Plant cell culture inherits most of the advantages of plant-based production systems, particularly the ability to produce complex proteins with correct posttranslational modifications without risk of contamination by human pathogens. Plant cell cultures also can be rapidly propagated in bioreactors as homogeneous suspensions for large-scale production, growing fast in simple synthetic media with doubling times as short as 16 h [5]. A breakthrough in plant cell culture technology was made in May 2012 by Protalix Biotherapeutics (http://www.protalix.com), an Israel biopharmaceutical company, for a plant cell-produced therapeutic enzyme — taliglucerase alfa — finally approved by the US Food and Drug Administration (FDA) as an orphan drug for Gaucher’s disease. Taliglucerase alfa is a hydrolytic lysosomal glucocerebrosidase for intravenous infusion and commercially known as Elelyso™; it is the world’s first plant-made human pharmaceutical and made by Protalix using carrot cells.

Plant cell lines most widely used for recombinant protein production are derived from tobacco (Nicotiana tabacum), particularly cultivar BY-2 (N. tabacum cv. Bright Yellow 2) cells. BY-2 cells are robust and fast growing and can multiply ≥100-fold in a week. They readily undergo Agrobacterium-mediated transformation and cell cycle synchronization [5, 81, 82]. Other commonly used cell lines include rice (Oryza sativa), alfalfa (Medicago sativa), and carrot (Daucus carota). In fact, these cell lines derived from common edible crops may be more favorable than tobacco cells in terms of by-product levels and regulatory compliance [81]. Rice cell suspension cultures are used almost as widely as tobacco BY-2 cells due to availability of the sugar-sensitive α-amylase promoter system (R Amy3D) [83]. This promoter is induced by sugar starvation and has enabled high-level expression of many pharmaceutical proteins in rice cells, e.g., α1-antitrypsin (rAAT) [33, 83], hGM-CSF [56], interleukin-12 [59], and human serum albumin [60], with highest secreted protein yields reaching 247 mg L−1 for rAAT [33]. While these production levels are impressive, growth rates, and general characteristics and stability of rice cell lines are inferior to those of tobacco BY-2 cell lines; rice cell viability is significantly decreased when cultivated in a sucrose-starvation medium [80].

The major bottleneck to full adoption of plant cell culture technology for commercial purposes has been low productivity with protein yields ranging from 0.01 to 10 mg L−1 [84]. To substantially improve protein expression for commercial success, strategies at both the molecular and at the process development levels are required to maximize efficiency of all stages of the production pipeline [5, 85]. This is similar to the strategy that has been systematically adopted by the mammalian cell culture industry over the past 25–30 years. Notably, in tobacco BY-2 cells, productivity of M12 mAb was ~8 pg cell−1 day−1 [86] vs. 20–40 pg cell−1 day−1 for Chinese hamster ovary (CHO) cells [79]. Besides low productivity, other major challenges remain including non-mammalian glycosylation, genetic instability, and cell culture scale-up in bioreactors [8, 34, 87].
3.1.2 Hairy Roots

Hairy roots are generated by infection of plant tissues with *Agrobacterium rhizogenes* that harbors a root-inducing (Ri) plasmid [88]. Similar to suspension cells, hairy roots can be grown in a controlled and sterile environment suitable for cGMP production of pharmaceutical proteins. However, as a more organized organ, hairy roots offer additional benefits, including genotype and phenotype stability and autotrophy for plant hormones [89, 90].

Hairy roots expressing a specific recombinant protein can be readily generated either by infecting stably transformed plants (expressing the target protein) with *A. rhizogenes* or by infecting wild-type plants with genetically modified *A. rhizogenes* harboring binary vectors containing the gene of interest [67]. After the first success producing a full-length murine IgG1 in tobacco hairy roots [91], more than 20 recombinant proteins, including reporter proteins (e.g., GUS and GFP), enzymes (e.g., human acetylcholinesterase and tPA), antibodies (e.g., human M12 mAb and murine 14D9 IgG1), antigens (e.g., HBsAg and cholera toxin B surface protective antigen), and cytokines and growth factors (e.g., interleukin-12, hGH, and hEGF), have been expressed in hairy roots [90, 92] with protein yields up to 3.3% TSP of an acetylcholinesterase [93] and >120 mg L\(^{-1}\) of GFP [68].

Hairy roots also secrete expressed proteins from cultured tissues, termed *rhizosecretion* [89, 94, 95], offering a simplified, low-cost approach for purification of foreign proteins from inexpensive and well-defined media. Because root tissue is not destroyed for recovery of secreted proteins, a given culture can be used for several cycles of bioproduction. Using an optimized induction protocol for the secretion of M12 mAb from tobacco hairy roots by addition of extra KNO\(_3\), α-naphthaleneacetic acid, and polyvinylpyrrolidone to the standard MS medium, antibody yield was improved by 30-fold, yielding 5.9 mg L\(^{-1}\) [62].

The bottleneck to exploiting hairy root technology for commercial purposes has been low protein productivity [90]. Expression systems have been strategically designed to include a strong promoter such as a double-enhanced *CaMV35S* promoter (2 × 35S) [93], a chimeric super-promoter (*Aocs, 3AmasPmas*) [96], and inducible promoters [97]. In addition, the special morphological characteristics of hairy roots including nonhomogeneous growth and highly branched phenotypes present major challenges to culture scale-up in bioreactors [98].

3.1.3 Moss

Moss protonema can be suspension cultured in bioreactors to provide another promising platform for producing recombinant products. While plant cells grow on sugar-based media without the need of light, moss is able to grow using light as a sole source of energy, needing only water and inorganic salts as a medium. This greatly reduces production cost and facilitates product recovery from the medium [51]. Moss also can perform complex posttranslational modifications of expressed proteins [99]. Because it relies on differentiated instead of undifferentiated plant cell cultures, moss cultures are genetically stable over long periods of time [100]. The moss *Physcomitrella patens*, with its genome fully sequenced in 2006 (http://www.cosmoss.org/), is the main species used for bioproduction.
A wide variety of biopharmaceutical human proteins has been produced in moss including tumor-directed mAbs with enhanced antibody-dependent cell-mediated cytotoxicity (ADCC) [101, 102], keratinocyte growth factor (FGF7/KGF) [77], asialo-erythropoietin (asialo-EPO) [73, 74], α-galactosidase and β-glucocerebrosidase, etc. [77] (www.greenovation.com). See a recent review [103]. Some of the recombinant biopharmaceuticals produced from moss are not only similar to those produced in mammalian cells, but even of superior quality (“biobetters”). For example, moss-made α-galactosidase lacks the terminal mannose phosphate and thus is taken up by cells via mannose receptors instead of mannose-6-phosphate receptors, yielding better pharmacokinetics in Fabry mice [103]. In addition, moss N-glycans are free of core α-1,6-fucose, a sugar structure typically present on the N-glycans of mammalian cell-derived proteins. The moss-made IgG lacking this sugar moiety was more efficient in antibody-dependent cell-mediated cytotoxicity (ADCC) than the mammalian cell-made counterparts [101–103].

A unique feature of *P. patens* is that its genome can be readily engineered through gene targeting, a gene replacement strategy based on homologous recombination [51]. The approach has been efficiently used for precision glyco-engineering of moss-produced proteins by knocking out or knocking in certain glycosyltransferase enzymes, allowing for production of humanized glycoproteins [101]. For example, moss mutants were engineered with genes encoding plant typical glycosyltransferases knocked out from the moss genome [72, 74] and further engineered to knock in the gene encoding β-1,4-galactosyltransferase into the xylosyltransferase or fucosyltransferase locus, respectively [104]. To avoid unwanted potential O-glycosylation at the hydroxyproline residues of human proteins, a gene responsible for prolyl hydroxylation was identified and deleted from the moss genome [99]. See recent reviews [105, 106].

### 3.1.4 Scaling Up In Vitro Systems

Large-scale cultivation of in vitro systems focuses mainly on cell suspensions and hairy roots. While there are many bioreactors that have been designed for in vitro culture as described in prior reviews that also cite specific examples (e.g., see [107–109]), only a few types of bioreactors seem to be sufficiently scalable for commercial application. These are briefly described below and schematically illustrated in Fig. 1.

While the stirred tank reactor (STR; Fig. 1a) is the industry workhorse for microbial systems, it is not the most ideal option for plant cultures, though as proof of concept some plant cell suspensions have been successfully scaled to 600 L [110, 111]. On the other hand, the bubble column, balloon, wave reactors, and variations thereof seem to be more effective thus far for scaling up in vitro cultures. Plant cell suspensions have been grown in nearly all types of liquid-phase bioreactors. The bubble column reactor (BCR; Fig. 1b) is easy to construct in-house with a variety of design variations. Gas enters at the base of the vertical cylinder through a frit forming small bubbles that rise through the chamber, resulting in aeration and mixing; gas vents through a sterile filter at the top. The BCR
unfortunately suffers from foaming. This led to development of the balloon-type bubble reactor (BTBR) (Fig. 1c) with a broader liquid surface area. This design alleviated foaming, providing better gas exchange than the BCR. While the BTBR scales from 4 to 500 L, it is constructed of glass, so at large scale it requires a stainless steel superstructure. This greatly increases capital costs and availability of these reactors is limited. The BTBR has been used to culture a wide variety of cell suspensions and hairy roots. See above cited reviews for more in-depth discussion of different requirements for growing cells vs. hairy roots including many species examples.

The wave bioreactor (Fig. 1d) is a horizontal, transparent plastic bag, seated on a slowly rocking platform that through wave action within the bag provides good agitation. With gas input into the headspace above the large surface area of the liquid in the bag, there is also good gas exchange. Although the wave reactor is being commercially used, it unfortunately scales horizontally with a relatively large
footprint compared to the vertical reactors, for example, that used by Protalix as subsequently discussed. Other than the glass balloon reactor, plastic single-use disposable culture chambers are also becoming the norm. These can include isolated bags or liners within a supporting superstructure. A vessel with a plastic liner was designed, demonstrated, and patented (US 6,709,862 B2) as versatile and functional some time ago by the Curtis lab [112]. For example, the SB200-X (OrbShake, Kühner AG) is comprised of a large 200 L cylindrical vessel with an inner disposable bag that sits on a platform and is orbitally shaken. This reactor has shown some success in cultivation of recombinant BY-2 tobacco cells for production of human M12 mAb. Yields of 300–387 g FW L\(^{-1}\) and ~20 mg L\(^{-1}\) M12 were equivalent to yields obtained in shake flasks [113]. Single-use cultivation chambers obviate cross contamination of products or cultures from run to run and are less capital intensive, and because there is no need for sterilize in place (SIP) or clean in place (CIP) plumbing, they also have simpler accommodation requirements [114]. Plastics approved for use by FDA include polypropylene, polystyrene, polyethylene, polytetrafluoroethylene, or ethylene vinyl acetate, facilitating regulatory approval of the overall production process [114].

Moss can also be grown in many of these same bioreactors [103]; however, if light is needed for autotrophic growth, then a photobioreactor is required along with attendant challenges that are further addressed in Sect. 3.2.3. Indeed, successfully grew \(P \). \(p\)atens under GMP-certified conditions in 100 and 500 L wave bioreactors under artificial illumination, so for high-value products smaller volume reactors may be reasonable. If moss could be grown heterotrophically, then the same bioreactors described for cells and hairy roots could be employed. Nevertheless, Greenovation Biotech GmbH (www.greenovation.com/) is using its BryoTechnology™ platform to commercialize recombinant biopharmaceuticals. Several therapeutic enzymes targeted for enzyme replacement therapies, such as \(\alpha\)-galactosidase, \(\beta\)-glucocerebrosidase, and complement factor H are under preclinical or Phase I development (www.greenovation.com/).

Modularity is a valuable tool in modern bioprocess design. Implementation of multiple, smaller scale, e.g., \(\leq 500 \) L, reactors is useful in production facilities because it provides the ability to scale in response to demand, minimizes risks of loss due to contamination, and is less capital intensive. An example of a successfully scaled up commercial cell suspension culture is that by Protalix using carrot cells engineered to produce recombinant glucocerebrosidase as a replacement therapy for managing Gaucher’s disease, taliglucerase alfa (Elelyso™) manufactured by Pfizer. The large-scale patented [115] cultivation system (ProCellEx™) for growing these carrot cells uses \(\geq 400 \) L vertical conical-shaped plastic culture bags similar in design and operation to a BCR with aeration inlet ports near the bottom of the bag and a gas vent/regulator at the top to help maintain bag inflation (Fig. 1e). Rising gas bubbles from the gas inlet ports provides culture agitation and mixing. The large suspension-filled bags have a grid-like superstructure providing external support (Fig. 1e). The system is modular, thereby allowing cultivation and harvest cycles; a central unit provides aeration and nutrients [116]. Inoculant and culture media are provided to
each reactor, and excess air and waste gases are removed. All units are housed in a clean room. Schematics and details of the reactor technology are in US Patent 2010/0112700 A1.

### 3.2 Aquatic Plants

Some aquatic plants also are promising bioproduction platforms, including duckweed and microalgae. Many functional industrial enzymes or human therapeutics have been expressed in duckweeds or microalgae at lower cost than in vitro systems [85].

#### 3.2.1 Duckweed

Duckweed, the common name for *Lemnaceae*, is a monocot plant family consisting of four major genera: *Lemna*, *Spirodela*, *Wolfia*, and *Wolffiella*. Duckweed is propagated clonally without the need for pollen or seeds, which simplifies line management, propagation, and the process feed stream [32, 117]. Duckweed is safe, fast growing in simple inorganic media (doubling time $\geq 36$ h), capable of making complex proteins, and easy to grow and harvest and has a high protein content (up to 45% dry weight) [85, 118]. Cultivation only requires inexpensive upstream facilities and is very scalable. The absence of pollen or seeds also makes duckweed environmentally safer than other transgenic flowering plants. Moreover, duckweed is edible, offering an attractive system for oral delivery [119, 120].

Duckweed can be transformed using either biolistics or *A. tumefaciens*. Efficient nuclear transformation protocols for two species of duckweed, *L. gibba* and *L. minor*, were established in 2001 [121], and $>20$ recombinant proteins were produced with expression levels up to 25% of TSP [32]. Products include industrial enzymes, e.g., E1 endoglucanase [122], and many therapeutic proteins, e.g., mAb [123], plasminogen [124], interferon $\alpha_2$ [125], vaccine antigen avian influenza including H5N1 hemagglutinin [126, 127], and M2e peptide [117].

In 1997, US-based Biolex, Inc. developed the duckweed (*Lemna*)-based expression (LEX) system and successfully expressed at least 12 therapeutic proteins, including small peptides, mAbs, and large multimeric enzymes [2]. However, in 2012, Biolex’s technology was sold to Synthon, a Netherlands-based pharmaceutical company, and interest in the LEX system subsided; research is currently focused on wastewater treatment (phytoremediation) and biofuel production.

#### 3.2.2 Microalgae

Microalgae integrates the merits of microbes, including rapid growth and ease of culture with those of higher plants in performing posttranslational modification and photosynthesis, thereby offering another promising platform for cost-effective production of recombinant proteins [128–130]. Microalgae have a very simple structure, grow in simple media, and produce large amounts of biomass with short life cycles [131]. Downstream purification of proteins from microalgae is similar to yeast and bacterial systems and thus is generally less expensive than from whole plants.
Many species also are generally regarded as safe (GRAS) for human consumption, thus providing a potential platform for oral delivery.

Both nuclear and chloroplast genomes of microalgae can be transformed for expressing heterologous proteins. However, due to the nuclear silencing, nuclear transformants generally accumulate less recombinant protein than chloroplast transformants [129]. Thus, the chloroplast expression system is currently regarded as more feasible for commercial production [128]. The disadvantage of chloroplast transformation is that this organelle lacks posttranslational modification capability, e.g., glycosylation [130]. However, this may provide a benefit for antibodies produced in chloroplasts, because glycan-free antibodies do not activate the immune system in humans [133, 134].

Chlamydomonas reinhardtii is the unicellular green algal species on which the majority of genetic engineering has been performed [128, 135, 136]. This alga grows fast with doubling time of ~10 h, supports easy nuclear and chloroplast transformation, and can be cultivated either photoautotrophically or with acetate as a carbon source [85]. A variety of high-value recombinant proteins, including antibodies, vaccines, growth factors, and industrial enzymes have been produced in microalgae from either nuclear or chloroplast transformation, as documented in several recent reviews [131, 137, 138]. Most of these were produced in the chloroplast of C. reinhardtii, but some were produced in other species, such as Dunaliella salina and Phaeodactylum tricornutum.

Development of economically viable bioproduction is still hampered by a lack of effective and consistent transformation methods for a wider variety of species, as well as low (nuclear expression) or inconsistent (chloroplast expression) recombinant protein yields [129]. PhycoBiologics (www.phycotransgenics.com) is currently attempting to commercialize microalgal production with indoor photobioreactors yielding axenic algae with >20% of TSP in the chloroplast (www.phycotransgenics.com) [132].

### 3.2.3 Scaling Up Aquatic Systems

Duckweed, moss, and microalgal aquatic species require light for autotrophic growth to achieve high biomass yields, which in turn affects product yields. Autotrophic cultivation requires either sunlight or significant artificial lighting intensity that is in close proximity to the cultured cells. Options include open systems such as ponds and raceways and closed systems such as tubular or plate photobioreactors. While possibly suitable for low-value commodity or industrial products, for products requiring strict GMP compliance, open systems are not recommended because they can be contaminated with a variety of other species and particulates and do not provide production source material consistency. There are also issues regarding environmental release of genetically modified species grown in an open pond. Although tubular or flat plate photobioreactors would seem best, these also have challenges. While the development of more cost-effective LED lighting is now available, these reactors cannot exceed certain culture depths. As culture density increases, light does not penetrate beyond several centimeters, and thus as the culture increases in density, growth decreases significantly. Although cGMP industrial
facilities now exist for moss cultivation, to our knowledge, there is none yet for microalgae [31]. For a more in-depth discussion of photobioreactor designs and limitations, see the review by Xu et al. [139].

### 3.3 Whole Plants: Stable and Transient Expression Systems

Whole plant expression still dominates plant-based bioproduction platforms and uses either transient expression via viral or nonviral vectors or stable transformation with transgenes targeted to either the nuclear or chloroplast genome. Stable transformation advantages include (1) a heritable transgene, permitting establishment of seed stock for future use, and (2) protein production scalable to field production. However, establishing and characterizing stable transgenic lines can be costly and time consuming. Other challenges include gene silencing, position effects, and GMO environmental concerns [132, 140].

When establishing stable transgenic plant lines, choices are often made regarding the gene integration site (nuclear vs. plastid), the subcellular compartment for recombinant protein accumulation (e.g., cytosol, apoplast, endoplasmic reticulum, and vacuole), and the plant tissue expression target (leaves vs. seeds). Decisions depend on a variety of considerations, including posttranslational modifications needed for protein functionality, stability of the expressed foreign protein in the plant host, desired expression levels of product, and downstream purification costs, as well as size and cost constraints of the product market [85, 95].

#### 3.3.1 Stable Expression: Leaf Based

Leaf tissues of nonfood crops have traditionally been used as a viable expression platform for either research or production purposes. The first plant-produced recombinant protein, an immunoglobulin, was expressed in transgenic tobacco leaves [141]. Tobacco has emerged as the leading plant species for leaf-based protein expression [142] because it produces high biomass yields (~300 tons per acre), is genetically well studied, and is readily amenable to genetic engineering [15]. Importantly, tobacco is a nonfood, nonfeed crop, which minimizes regulatory barriers by eliminating the risk of plant-made recombinant proteins entering the food supply [143]. The first clinical trial of a plant-produced biopharmaceutical was the secretory antibody variant of Guy’s 13 produced in field-grown tobacco leaves by Planet Biotechnology Inc. [144]. Besides tobacco, other leafy crops, e.g., lettuce, alfalfa, and clover, have also been exploited as an expression platform. Alfalfa is advantageous because it is a perennial that fixes nitrogen and displays notable homogeneity of N-glycosylated recombinant proteins [85, 145].

Both leaf nuclear and plastid genomes have been targeted for integration of heterologous genes. The choice of gene integration site is generally dictated by the posttranslational requirements of the target proteins. Nuclear integration is necessary for expression of functional glycoproteins, such as EPO and tPA, that need proper processing in the endo-membrane system [35]. Thus far >100 functional proteins have been successfully expressed in leaves with nuclear transformation. These
include mAbs [141, 146–148], vaccines [149], cytokines [150, 151], and industrial enzymes [152, 153]. See reviews [3, 84, 132, 154]. Unfortunately, low copy of gene integration, gene silencing, and proteolytic degradation in the aqueous environment of leaves resulted in <1% TSP [155, 156].

Each plant cell contains an average of 50–100 chloroplasts, and each chloroplast contains ~100 copies of its genome, thus the chloroplast enables thousands of copies of a given transgene to be expressed [157–159]. With chloroplast transformation, accumulation of recombinant protein up to 70% of the total leaf protein has been achieved [160]. See the special issue on Chloroplast Biotechnology in Plant Biotechnology Journal (June 2011). A wide range of proteins, ranging from very small antimicrobial peptides or hormones to very large viral or human proteins, have been successfully expressed in plant chloroplasts [161]. Except for exceptionally high-level expression, therapeutic proteins accumulated in the chloroplasts of lyophilized plant cells can be stored for several months or years without a decrease in their functionality, thereby eliminating costs of cold storage and transportation [162]. However, because the chloroplast does not support many important posttranslational modifications desired for expressing complex proteins, this technology is limited to production of proteins whose functions are independent of glycosylation. Another disadvantage of chloroplast transformation is the difficulty of transforming many important crop plants using current methods [163].

Chloroplast transformation also provides a unique bioproduction system for oral medicines [164]. When a pharmaceutical protein is accumulated in plant leaves, the plant cell wall can protect the protein drug from acids and enzymes in the stomach via bioencapsulation after oral delivery. However, microbes residing in the human gut have evolved to break down the plant cell walls to release the target protein [164]. If a transmucosal carrier (receptor-binding protein) such as cholera toxin subunit B (CTB) and heat-labile toxin subunit B (LTB) that binds to GM1 receptors is fused to the protein drug, it will efficiently cross the intestinal epithelium and be delivered to the circulatory or immune system [161]. Several drugs have advanced to the clinic, including glucocerebrosidase for treating Gaucher’s disease [12], clotting factor IX for treating hemophilia B [165], myelin basic protein for treating Alzheimer’s disease [160], acid α-glucosidase for treating Pompe disease [166], and, most importantly, a variety of oral vaccines for treating infectious diseases [158, 167]. See recent reviews [164, 168, 169].

Limitations of leaf-based platforms compared to seeds are the short shelf life of leafy tissue and the high variability of the production system. Plant leaves degrade faster upon harvest and must be processed immediately to ensure product stability and quality. Product yields in field-grown materials can be highly variable due to environmental changes (both biotic and abiotic). Finally, large-scale, regulatory-compliant disposal of transgenic biomass waste may have volume and cost implications [85].

### 3.3.2 Stable Expression: Seed Based

Recombinant protein expression targeted to plant seeds can overcome the major limitations associated with leaf tissue: protein stability and storage [170, 171]. Plant
seeds are naturally suited to produce and store proteins. High protein content (7–10%), low protease activity, and low water content make seeds an attractive alternative bioproduction platform [155, 156]. Antibodies or vaccines expressed in cereal seeds remained stable at ambient temperatures for years [85, 172]. In addition, edible seeds such as maize and rice have GRAS status making them particularly suitable for developing oral vaccines that can be administered as flakes or flour with minimal purification [173]. In the past decade, significant advances have occurred using seeds as bioreactors including the commercialization of the first plant-derived recombinant protein, avidin, produced in maize seeds [170].

Cereal seeds, e.g., rice, wheat, barley, soybean, and maize, are commonly used as expression hosts [174]. Other typical production platform seed crops include legumes (pea, cowpea, and soybean), oil crops (canola and safflower), tobacco, and Arabidopsis. Stable expression yields up to 10% of total seed proteins have been achieved [170, 175]. Products include therapeutic proteins such as antibodies [176], vaccines [177], and cytokines [178]; industrial enzymes such as trypsin [179], phytase [180], and cellulase [181]; and biopolymers such as spider silk protein [182]. See recent review [173].

Maize seeds are the most widely used host for commercial therapeutic proteins and industrial enzymes [173]. Compared to other cereals, maize has a larger grain size, a higher proportion of endosperm (≥82% of the seed), and a higher biomass yield per hectare at lower production costs [173, 183, 184]. Many industrial enzymes, including GUS, cellulase, laccase, and trypsin, have been successfully produced in maize seeds and marketed [27, 184]. Therapeutic proteins such as the HIV neutralizing antibody 2G12 [185], influenza virus H3N2 nucleoprotein [186], and α-galactosidase [187] in maize seeds were produced cost-effectively with simple downstream purification processes. Other seed-based platforms in commercial development include rice seed-produced human transferrin [188] by US-based Ventria Biosciences (www.ventria.com/) and barley grain-produced endotoxin-free growth factors and cytokines (Orfeus™ expression system) [189] by Iceland-based ORF Genetics Ltd. (www.orfgenetics.com/).

Seeds from some oil crops such as safflower and rapeseed have been used to develop a novel “oleosin-fusion” bioproduction platform, in which recombinant proteins are expressed as a fusion with oleosin, an endogenous protein that localizes within oil bodies of rapeseed and safflower [190, 191]. Fusion proteins accumulated in the oil bodies are easily separated in the lipid fraction from the bulk seed homogenate. SemBioSys Genetics, a Canadian biotech company, was developing and commercializing this technology for the low-cost “biosimilar” insulin [85], but operations ceased in 2012 due to financial problems.

Although a seed-based platform has many positive attributes, major hurdles still exist. Compared to the leaf-based platform, seeds have a relatively lower biomass and high possibility of gene leakage into the environment via the seed or from pollen dispersal [15, 192]. There is strong reluctance among scientists, regulators, and the general public to use seeds of major crops (e.g., maize, rice, and wheat) for recombinant protein production because of the possibility of contaminating the food chain [192]. Considerable time is also required to create high-expressing
transgenic plant lines [155]. However, the stability of foreign proteins, post-harvest processing, and overall cost makes the seed-based platform suitable for many recombinant proteins produced on a large scale [193].

3.3.3 Transient Expression

Transient expression is achieved either by epichromosomal expression of *A. tumefaciens* directly infiltrated into plant tissues (agroinfiltration) or by viral-based expression vectors [194, 195]. During transient expression, foreign genes are typically introduced into leaves of plants by vacuum infiltration of *Agrobacterium* containing genes of interest in an expression vector [85]. Recombinant protein production within the plant tissues (usually leaves) can be initiated quickly, within 24 h, continuing for several days depending on vector and target protein. Because a high copy number of foreign genes is introduced into each plant cell and free of “position effect” on gene transcription, the protein yields are usually higher in transient than in stably expressed plants [196]. Since there is no transgenic plant created, this also addresses regulatory issues and public concerns for GMO plants [197]. As a result, transient expression has been increasingly used for production of biopharmaceuticals, in particular, antigen vaccines and antibodies [195, 197].

*N. benthamiana* is the most common host plant for transient expression because it is amenable to genetic transformation and rapidly yields large amounts of biomass from seeds for scale-up production [198]. Potato, green pea, *Arabidopsis*, lettuce, and other *Nicotiana* species (e.g., *N. debneyi*, *N. excelsior*, and *N. simulans*) also serve as alternative hosts for transient expression [85]. There are two basic strategies for introducing transgenes into host plant cells for bioproduction: nonviral vector based or viral mediated [199, 200]. Nonviral vector-based expression using common plant expression vectors enables rapid accumulation of recombinant proteins in plant leaves, typically 2–4 days post-infiltration, with protein yields of 0.1–200 μg g⁻¹ FW [85]. The viral-mediated expression process takes ~2 weeks for protein expression, but generally produces higher levels of recombinant protein up to 5.0 mg g⁻¹ FW GFP [201]. Some effective viral vector-based expression platforms include Geneware® (Kentucky BioProcessing LLC) and magnICON® (Icon Genetics) based on a tobacco mosaic virus (TMV) RNA replicon and the geminiviral expression system based on a bean yellow dwarf virus (BeYDV) DNA replicon (Arizona State University) [202–204].

Transient systems also provide the ability to simultaneously co-express several genes to produce complex proteins, such as antibodies, viruslike particles (VLP), and other multichain proteins [195]. Numerous therapeutic proteins, especially those addressing sudden viral epidemics (e.g., an outbreak of Ebola, severe acute respiratory syndrome, or influenza pandemic) have been successfully produced by a transient expression system [132, 205, 206]. For example, Mapp Biopharmaceutical Inc. (San Diego, CA) used the geminivirus technology to transiently express the humanized antibodies MB-003 (Mapp) and ZMab in tobacco leaves (http://mappbio.com/z-mapp/). An optimized cocktail combining the best components of the MB-003 and ZMab was ZMapp™, which cured 100% of Ebola-infected rhesus macaques primates [207, 208]. In another example, plant-produced influenza
vaccine candidates, such as VLP antigens against avian flu (H5N1) virus and swine flu (H1N1) virus, were developed by the Canadian biotech company Medicago Inc. (Quebec, Canada) and have undergone Phase I and II clinical trials with positive results (http://medicago.com/pandemic-flu/). Plant-produced influenza vaccines are regarded as quicker to develop and potentially cheaper than egg-produced vaccines. Many other therapeutic proteins reportedly produced by the transient expression platform include IgG and IgA antibodies [209, 210], vaccine antigens against malaria, influenza and HIV [211–215], and therapeutic enzymes treating lysosomal storage diseases [216]; see recent reviews [140, 195, 217, 218].

Transient expression is easily applied to industrial scale for mass production of recombinant proteins. Milligram to gram quantities of target proteins can be rapidly produced in weeks to allow animal and clinical testing. Several plant biotech companies, including Kentucky BioProcessing LLC (Owensboro, KY), Medicago Inc. (Quebec, Canada), Fraunhofer CMB (Newark, DE), Mapp Biopharmaceutical Inc. (San Diego, CA), and iBio Biotherapeutics (Bryan, TX), have developed cGMP manufacturing facilities for biopharmaceuticals using vacuum agroinfiltration of N. benthamiana [219]. A major disadvantage of the transient system is the necessity of harvesting at a particular time. If using the product in humans or animals, an additional purification step to remove endotoxins derived from the infiltrated Agrobacterium is required [85]. Transient expression systems have been continuously optimized in the past decade for rapid, high-yield, and large-scale production of recombinant proteins, which expedites the acceptance of this production platform for the commercial production of a broad range of biopharmaceuticals.

### 3.3.4 Scaling Up Whole Plant Systems

While field production of plants provides a distinct advantage given its apparent low cost, considerations of weather, pests, and seasonal effects on productivity will increase costs. Similarly there are concerns regarding the risk of unintended environmental contamination of native gene pools by genetically modified field crops. An alternative is the use of controlled environment agriculture (CEA), where the transgenic crop is grown in glasshouses. Of course this increases production cost substantially, but may be worthwhile if there is high product value, e.g., for a therapeutic. CEA is not subject to seasonal variation, providing greater control over cultivation conditions and thus the quality of the crop and its contents. A number of companies have developed more or less automated cultivation systems under glass where product quality is maintained and aseptically handled in its final form. Medicago has developed such a CEA system whereby they hope to produce a number of different products via transient expression in tobacco. The production system begins after there is a synthesized gene of interest (GOI) from, for example, a virus. Then young N. benthamiana tobacco plants grown in pots in contained greenhouses are handled robotically including the vacuum infiltration step. Large pallets of plants are inverted into a solution containing the GOI and vacuum infiltrated. Afterward the inverted plants are drained and rotated back into vertical position and then moved into another more stringent culture facility for a 4–6-day
incubation for gene expression and protein synthesis. Plants are subsequently moved
to a harvesting area where leaves are stripped by hand and then extracted. Using
methods applicable to any other clinical grade in vitro or microbial product, the
target protein is purified. The process is documented in this video: https://www.
youtube.com/watch?v=lAk_HkFi9-s.

There are different GMP concerns for products produced from whole plants that
extend beyond those for products produced from in vitro cultures (Fischer et al.
2012). Guidelines drafted in the USA by USDA and FDA exist to cover all platforms
described herein (FDA-USDA 2002 http://www.gmp-compliance.org/guidemgr/
files/BIOPLANT.PDF ). These include selection of a crop species platform suitable
for industrial production; in industry a diversity of platforms is problematic because
of the need for consistent and stringent regulatory compliance, especially for ther-
aputics. Subsequently there should be an early stage determination of field vs. CEA
cultivation. Similar to in vitro or microbial production, master seed or tissue banks
must be established and maintained to provide batch to batch consistency. Isolation
of GM plant material, controlled seed stock, APHIS/BRS permits for field-grown
plants, appropriate confinement for transport of source plant material from field/
greenhouse to production facility, control of harvested material and its transfer to the
processing facility, waste biomass control, storage control, are all needed for pro-
duction operations that can be validated.

Whole plant production can be further streamlined through second generation
propagation via shoot regrowth from a retained axillary bud on the rootstock after
harvesting the initial shoot Kim [220]. Functionality and yields of protein in
harvested leaves from three successive harvests remained relatively consistent.
Time required for seed production and germination and second and third generation
shoots required >50% less time to harvest vs. initial planting. Another obvious
alternative strategy would be rooting of clonal cuttings of transgenic plants to rapidly
scale-up whole plants. Such a strategy should also minimize time to harvest as there
is no need to generate seed or use in vitro micropropagules.

4 Commercialization Status and Outlook

Since conceptualization in 1989, the plant molecular farming industry has grown
rapidly. While some plant-made industrial proteins/enzymes (e.g., avidin, GUS, and
trypsin) and one plant cell-made pharmaceutical for human use (Eleyso™) have
been commercialized, many are in various developmental stages. Examples of some
successful plant-produced products, either on the market or in commercial develop-
ment, are listed on the Molecular Farming website last updated in March 2014
(http://www.molecularfarming.com/PMPs-and-PMIPs.html), or in some recent
reviews [50, 132, 221, 222]. The major biotech companies involved in research
and development of plant-produced protein products that are still in business are
listed in Table 3. These companies include the representatives of the very first
commercial ventures (Large Scale Biology Corp., Planet Biotechnology, Ventria
| Company name and country            | Plant species/ expression platform | Main products                                                                 | Status of products | Status of companies                                  |
|------------------------------------|------------------------------------|-------------------------------------------------------------------------------|--------------------|-----------------------------------------------------|
| Biolex Therapeutics, USA           | Duckweed/stable expression – LEX platform | Anti-CD20 antibody, controlled release interferon α2b for the treatment of HCV and HBV, etc. | Preclinical and Phase I, II, and III clinical trials | LEX platform sold to Synthon in 2012                 |
| Epicyte Pharmaceutical, USA        | Maize, tobacco/seed-based and leaf-based expression | Antibodies for infectious and inflammatory disease | Preclinical development | Acquired by Biolex in 2004                           |
| Fraunhofer IME, USA, Germany       | Tobacco/leaf based                    | HIV antibody                                                                   | Phase I clinical trials | Active                                               |
| Fraunhofer CMB, Germany            | N. benthamiana/ transient expression | Influenza vaccine                                                              | Phase I clinical trials | Active                                               |
| Greenovation Biotech GmbH, Germany | Moss (P. patens)/ stable expression – BryoTechnology™ platform | α-Galactosidase, β-glucocerebrosidase, and factor H                             | Preclinical and Phase I clinical trials | Active                                               |
| Kentucky BioProcessing Inc., USA   | N. benthamiana/ transient expression – Geneware® platform | Aprotinin and pharma (antibody ZMapp™, vaccines, and therapeutic enzymes)       | Aprotinin marketed by Sigma-Aldrich. Others under Phase I and II clinical trials | KBP (active now) offers contract biomanufacturing services; LSBC out of business in 2006 |
| iBio Biotherapeutics, USA          | N. benthamiana/ transient expression | iBio-CFB03 for anti-fibrosis                                                    | iBio-CFB03 granted orphan drug designation in 2016 | Active                                               |
| Icon Genetics, Germany             | Tobacco/transient expression – magnICON platform | Vaccine for non-Hodgkin’s lymphoma                                              | Phase I clinical trial | Purchased by Nomad Bioscience in 2012               |
| Mapp Biopharmaceutical, USA       | Tobacco/transient expression – magnICON platform | Antibody (ZMapp™)                                                              | Phase I and II clinical trials | Active                                               |
| Company                        | Plant/Expression System                              | Product/Application                                      | Status                                      | Acquirer (Date)                      |
|-------------------------------|-----------------------------------------------------|-----------------------------------------------------------|---------------------------------------------|--------------------------------------|
| Medicago, Canada              | Alfalfa, *N. benthamiana*/ transient expression — Proficia™ platform | Pharma (vaccines for influenza, rabies, and rotavirus)   | Phase I and II clinical trials             | Acquired by Mitsubishi Tanabe Pharma in 2013 |
| Nomad Bioscience, Germany     | Tobacco/transient expression — magniICON platform   | Cellulase enzymes and antimicrobial proteins (colicins, lysins, and bacteriocins) | Development                                | Active                               |
| ORF Genetics Ltd., Iceland    | Barley/seed-based expression                        | Growth factors and cytokines                              | Marketed for diagnostics, research, and cosmetics | Active                               |
| Phycobiologics Inc., USA      | *Chlamydomonas*/ algal-based expression             | Vaccine growth factor and enzymes                         | Under development                           | Active                               |
| Planet Biotechnology, USA     | Tobacco/leaf-based expression                       | CaroRx™ for dental caries; PBI-220                        | CaroRx™ under Phase II clinical trial; PBI-220 Phase I clinical trial | Active                               |
| Protalix Biotherapeutics, Inc., Israel | Tobacco or carrot/cell suspension culture — ProCellEx® platform | Pharma (β-glucocerebrosidase, α-galactosidase, DNase I, anti-TNFα) | One commercialized; others under clinical trials (Phases I/II/III) | Active                               |
| Synthon, Netherlands          | Duckweed/stable expression — LEX platform           | Anti-CD20 antibody for non-Hodgkin’s lymphoma and thrombolytic drug | Preclinical development                     | Active                               |
| Ventria Bioscience, USA       | Rice/seed-based expression — ExpressTec platform    | Lactoferrin and lysozyme; VEN120 for inflammatory bowel disease; VEN150 for HIV-associated chronic inflammation; nontherapeutics for basic research | Therapeutics under Phase II clinical trial or clinical development; nontherapeutic products marketed | Active                               |
Bioscience, SemBioSys Genetics, and Protalix), companies established in the mid-1990s (Epicyte, Medicago, Biolex, Icon Genetics, Greenovation, and ORF Genetics), and companies that emerged from the earlier pioneers (Mapp Biopharmaceutical, Kentucky BioProcessing, and Nomad Bioscience) [219]. In 1999, Large Scale Biology Corporation (Owensboro, KY), now Kentucky BioProcessing, designed and opened the first cGMP manufacturing facility for production of recombinant therapeutics by using the plant-virus transient expression system Geneware®. For plant transient expression systems (vacuum infiltration of *N. benthamiana*) to rapidly manufacture vaccines in response to epidemics and outbreaks, the US Defense Advanced Research Projects Agency (DARPA) sponsored four pilot-scale cGMP manufacturing facilities in the USA, including Fraunhofer CMB (Newark, DE), Kentucky BioProcessing (Owensboro, KY), Medicago (Durham, NC), and Caliber Biotherapeutics (now iBio Biotherapeutics, Bryan, TX) [206]. So far, iBio Biotherapeutics runs the world’s largest plant-based vaccine production facility with the capacity to process over 3500 kg of plant biomass per week (http://www.ibioinc.com/). With the advance in both technologies and manufacturing facilities, plant molecular farming is now reaching the stage at which it may challenge established microbial and mammalian bioproduction systems. Commercialization status of the three classes of plant-made recombinant products is briefly described as follows.

4.1 Plant-Produced Industrial Proteins/Enzymes

Due to ProdiGene’s pioneering work on the development and commercialization of several hydrolases including avidin, GUS, lysozyme, trypsin (TrypZean™), and aprotonin (AproliZean™) in the late 1990s [3, 7, 223, 224], plant-made industrial proteins spearheaded commercialization over the other two classes of proteins [85]. However, ProdiGene went bankrupt in 2002 due to mishandling US field tests of maize genetically modified to make pharmaceutical products and received more than a $250,000 fine from USDA. Since then, efforts on commercializing plant-made industrial proteins have declined. Most of the research is now focused on production of biomass-degrading enzymes for the biofuel industry, e.g., cellulase, hemicellulase, xylanase, ligninase, α-amylase, and laccase [181, 225–227]. For example, an Arkansas-based start-up company, Infinite Enzymes (http://www.infiniteenzymes.com/), has been developing a cost-effective maize seed production system for cellulase enzymes for cellullosic biofuel production. It is noteworthy that in 2011, a transgenic variety of maize expressing thermostable α-amylase was approved by the USDA for commercial use as an improved biofuel feedstock. This transgenic maize line, marketed as Enogen™, was developed by Syngenta (http://www2.syngenta.com/en/index.html) to produce — within the kernel — an enzyme needed to break down starch for biofuel production. However, it sparked a controversy not only from anti-GMO organizations but also from some biotech supporters because of environmental and human health issues [228].
4.2 Plant-Produced Therapeutic Proteins

This class of protein products has attracted the most interest in research and development in molecular farming. More than 20 plant-made pharmaceuticals have been placed in preclinical or clinical trials to date [132], and specific product examples are further discussed.

4.2.1 Antibodies

Production of therapeutic antibodies (plantibodies) is of great interest for the plant molecular farming industry [209]. Although none to date has been approved by pharmaceutical regulators, several plant-made antibodies have made it to human clinical trials. For example, CaroRx™, a secretory IgA (SIgA) plantibody produced by Planet Biotechnology Inc., was the world’s first clinically tested antibody for preventing adhesion of decay-causing bacteria to the tooth surface. Since 1999 CaroRx™ was subjected to Phase II clinical trials (topical) for dental caries in the USA under an FDA-approved Investigational New Drug Application [229], but was discontinued in early 2016. ZMapp™, a cocktail of three mAbs produced in tobacco leaves by Mapp Biopharmaceutical Inc. to combat the 2014 Ebola virus outbreak in Africa, underwent clinical Phase I and II trials in 2015 in the USA, Liberia, Sierra Leone, and Guinea, sponsored by the National Institute of Allergy and Infectious Diseases (NIAID) [132]. In September 2015, ZMapp™ was granted a fast-track status by the US FDA (http://mappbio.com). Although it has not yet received final approval by FDA, to date, ZMapp™ is the only drug that has been effectively used to treat patients infected with the Ebola virus [206]. In addition, a tobacco-derived HIV neutralizing antibody 2G12, produced by an EU funded project, Pharma Planta, recently completed a Phase I clinical trial [147].

4.2.2 Vaccine

Although several plant vaccines, for either animals or human, are now in clinical trials with encouraging results [229, 230], none thus far has been commercialized. Despite Dow AgroSciences receiving the world’s first regulatory approval by USDA in 2006 for a tobacco cell-produced vaccine against Newcastle disease virus in chickens, it only remained a proof of concept; Dow AgroSciences never intended to market the product [231]. However, since the scale of animal vaccination is so large, plant-based systems may represent the only cost-effective production platform on a scale for which other (non-plant-based production) methods are not competitive. Furthermore, plant-made veterinary vaccines, such as those made in seeds, fruits, and leaves, can be orally delivered as part of the animal feed, thus offering great convenience and economy in immunizing large populations of animals on farms [232]. In terms of commercialization potential, regulatory approval for the plant-made veterinary vaccines can be significantly less onerous than that for human vaccines [233]. Therefore, the most likely near-term possibilities for commercialization of plant-derived vaccines will be the veterinary products. Key examples of plant-produced vaccines tested in target animal species, including those against avian influenza, foot-and-mouth disease, Newcastle disease, diarrheal disease caused
by enterotoxigenic *E. coli*, porcine reproductive and respiratory syndrome, and swine transmissible gastroenteritis, are listed in recent reviews [231, 234]. These antigens were expressed in potato tubers, rice seeds, corn seeds, and *N. benthamiana* leaves. With continuing efforts to optimize the bioproduction platform, successful commercialization of plant-made veterinary vaccines is imminent.

The recent outbreak of avian and swine flu spurred development of seasonal and pandemic influenza vaccines. Medicago developed the Proficia™ technology, a transient plant expression platform for rapid and high-yield production of vaccines and antibodies, in particular, the viruslike particle (VLP)-based antigen (VLPExpress™ platform). Their products are at various stages of development. For example, the vaccine candidates against various influenza strains (e.g., H5N1 and H1N1) in Phase I and II human clinical trials were found safe and well tolerated, and potency was among the most effective of the industry (www.medicago.com). Mitsubishi Tanabe Pharma (Osaka, Japan) acquired Medicago in 2013 and announced that its tobacco-based flu vaccine could hit the market by 2018 or 2019. Fraunhofer CMB (Newark, DE) developed another type of plant-made VLP vaccine, Pfs25-VLP, for blocking malaria transmission [236]. The Phase I trial of Pfs25-VLP was completed in 2016 (https://clinicaltrials.gov/ct2/show/NCT01867463). One of the unique features of plant-made vaccines is that plants not only serve as the production “bioreactor” but can serve as the delivery vehicle for oral vaccines [236, 237]. Professor Charles J. Arntzen, a pioneer in plant-made oral vaccines at Arizona State University, has been developing plant-based oral vaccines including HBsAg (Phase I in lettuce and Phase II in potato), *Vibrio cholerae* vaccine, heat-labile toxin B subunit of *E. coli*, and the capsid protein of Norwalk virus (all are Phase I in potato) [1, 2, 85].

4.2.3 Therapeutic Enzymes
Therapeutics targeted for enzyme replacement therapies have been mainly developed by the Israel biopharmaceutical company Protalix Biotherapeutics. They developed and commercialized several recombinant therapeutic proteins through their ProCellEx® plant cell-based expression system (www.protalix.com). In 2012 Protalix commercialized Elelyso™ (taliglucerase alfa for injection), the very first plant-made therapeutic ever approved by the FDA for human use and has been marketed in the USA, Canada, and many Latin American countries (Uplyso™ in Latin America). Elelyso™ has comparable enzymatic activity and uptake in macrophages and is structurally homologous to Cerezyme® (imiglucerase), manufactured in CHO cells by Genzyme [57, 238]. Elelyso™, however, is ~25% less expensive than Cerezyme® because the plant cell-synthesized enzyme does not require further modification of its N-glycans for clinical use after bioproduction (terminal mannose residues are already exposed) [57]. Protalix’s development pipeline for therapeutic enzymes includes PRX-102, a pegylated version of a recombinant human α-galactosidase enzyme for the treatment of Fabry disease; PRX-110, a chemically modified DNase I for the treatment of cystic fibrosis; and PRX-112, an orally delivered glucocerebrosidase enzyme for treatment of Gaucher’s disease. To date,
Protalix has successfully completed a Phase I clinical trial of PRX-112 in Gaucher’s patients. In June 2016, the company initiated its Phase III clinical trial of PRX-102.

There are also three moss-derived enzyme products targeted for enzyme replacement therapies: α-galactosidase for Fabry disease, β-glucocerebrosidase for Gaucher’s disease, and factor H for atypical hemolytic uremic syndrome are under Phase I and/or preclinical development by the German biotech company, Greenovation (www.greenovation.com/). With the cGMP-compliant manufacturing process well established in Greenovation, the first moss-expressed biopharmaceutical is expected to come to market in the near future.

4.2.4 Other Therapeutics

Other plant-made therapeutics such as human growth factor and cytokines are in clinical trials or are already on the market. For example, using barley grain as a production host (Orfeus™ platform), ORF Genetics produced high-grade, animal-free, and endotoxin-free human growth factors and cytokines (ISOkine™ growth factors and cytokines) targeted for stem cell research (www.orfgenetics.com/). About twenty ISOkine™ products, such as IL-3, GM-CSF, EGF, TNF, SCF, etc., are also on the market (http://orfgenetics.com/ISOkine/ProductList/). Commercialization of several moss-made human growth factors, including FGF7/KGF (keratinocyte growth factor), EGF, and HGF (hepatocyte growth factor) intended for mammalian cell culture, has also been achieved by Greenovation [77]. FGF7/KGF is the first commercially available moss-made human protein. In addition, PRX-106, an oral formulation of anti-TNFα for treatment of immune and inflammatory-mediated disorders, is through Phase I clinical trial by Protalix, showing that the drug was safe and well tolerated and had gut biological activity and induction of regulatory T cells (www.protalix.com).

4.3 Plant-Produced Biopolymers

Compared to industrial enzymes and therapeutic proteins, commercial development of plant-made protein biopolymers lags. The major biopolymer products expressed in transgenic plants include collagens [239], spider silk proteins [22, 182], elastin-like polypeptides [15], and plant gum [240], which are promising biomaterials for regenerative medicine and tissue engineering as well as for the food industry [241].

5 Conclusions

Large-scale production and commercialization, especially of therapeutic proteins using plant production platforms, has passed proof of concept. While there are always challenges for each product and downstream process efficiency can certainly be improved, the technology for producing recombinant proteins in plants is past its infancy. More work is still needed to realize the full potential for plant-made industrial proteins and enzymes. Although not all industrial sectors have equally
progressed, the list of successfully commercialized plant-produced recombinant proteins is rapidly growing, suggesting a bright future for the biotechnology industry.

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