The MultiBac Baculovirus / Insect Cell Expression Vector System for Producing Complex Protein Biologics

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ABSTRACT

Multiprotein complexes regulate most if not all cellular functions. Elucidating the structure and function of these complex cellular machines is essential to understanding biology. Moreover, multiprotein complexes by themselves constitute powerful reagents as biologics for the prevention and treatment of human diseases. Recombinant production by the baculovirus / insect cell expression system is particularly useful for expressing proteins of eukaryotic origin and their complexes. MultiBac, an advanced baculovirus / insect cell system has been widely adopted in the last decade to produce multiprotein complexes with many subunits that were hitherto inaccessible, for academic and industrial research and development. The MultiBac system, its development and numerous applications are presented. Future opportunities for utilizing MultiBac to catalyze discovery are outlined.
1/ Introduction –The baculovirus expression vector system (BEVS)

More than 30 years ago, the high level production of a heterologous protein by using an insect specific baculovirus, derived from the Autographa californica multiple nuclear oolyhedrosis virus (AcMNPV) was reported. Max Summers and co-workers produced functional human IFN-β in insect cells infected by a recombinant baculovirus [1]. This development was made possible by the previous observations that late in its viral life cycle, baculovirus expresses at very high levels a protein, polyhedrin, which is not essential in laboratory culture. Substitution of the polyhedrin gene in the baculoviral polh locus by a foreign gene of interest resulted in comparably high-level expression of the desired gene product, driven by the polh promoter, without compromising virus infectivity and the viral life cycle [2]. Shortly after, a second study by Lois Miller and colleagues demonstrated that another very late promoter p10, showed similar characteristics and could also be used for high-level production of heterologous proteins [3].

These two seminal studies established the baculovirus / insect cell expression system as a powerful means to produce proteins recombinantly. In the three decades since these hallmark contributions, baculovirus expression has become a widely adopted technology for academic and industrial applications, in research and development as well as manufacturing, and a wide range of proteins have been made by baculovirus expression vector systems (BEVS) [2, 4-7]. Multicomponent virus like particles (VLPs) resembling complex virus shells, have been successfully produced with BEVS, including VLPs from bluetongue, rotavirus and others [8-11]. More recently, the first baculovirus produced proteins have been approved in the therapy or prevention of human disease, including vaccines against influenza (Flublok®) and cervical cancer (Cervarix®), and immune-therapeutics against tumors of the prostate (Provenge®) [4]. Moreover, baculovirus itself has emerged as a versatile tool for
gene therapy, either as a production system for recombinant adeno-associated viruses [12-14] or as a DNA-based gene delivery vehicle in its own right [15,16].

The development of BEVS for heterologous protein production and its manifold exploits have been authoritatively reviewed recently in a number of contributions, recapitulating comprehensively the technical aspects of this technology [2, 4, 5]. The subject of this present contribution is MultiBac, a particular baculovirus expression vector system developed and implemented more recently [17-25]. MultiBac was originally conceived to meet the imposing challenge of producing eukaryotic multiprotein complexes, vital cornerstones of biological activity, in high quality and quantity for high-resolution structural and functional analysis. The system has been uniquely successful in catalyzing multiprotein complex research globally. MultiBac, its ongoing development, its numerous applications and future prospects are reviewed in the following.
2/ The MultiBac system for expressing eukaryotic multiprotein complexes

Protein complexes catalyze key functions in the cell, and as a consequence, are an intense focus of contemporary biological research efforts. Genomics and proteomics studies have underpinned that most if not all proteins in eukaryotic cells are parts of larger assemblies, with in humans often comprise ten or more individual subunits. The complex interplay of proteins in these complexes is essential for maintaining cell homeostasis and biological activity, and for development. High-resolution functional and structural characterization of the large number of multiprotein assemblies in the cell is critical to understanding cell biology [20, 26, 27].

Multisubunit complexes may be purified from their native cell environment and their structure and function analyzed successfully at near-atomic resolution, provided they are sufficiently abundant and homogeneous. Well-known examples include RNA polymerases and ribosomes [28-31]. The overwhelming majority of multiprotein complexes in the cell, however, are characterized by low or very low abundance, which considerably complicates or even rules out their purification from native source material. Furthermore, it is becoming increasingly clear that many proteins may exist not only in one, but a number of distinct complexes, carrying out diverse functions depending on the partner molecules they associate with at a given time. Together, this often obstructs obtaining compositionally pure and homogenous material by classical fractionation of cells and subsequent biochemical purification, notwithstanding significant progress notably in endogenous tagging methods to genomically modify endogenous proteins by powerful extraction aids such as tandem affinity purification (TAP) tags, for instance [32-34]. A solution to these issues is recombinant overproduction, enabled by the development and implementation of powerful overexpression technologies that can achieve high level production of homogeneous and active eukaryotic complexes for detailed mechanistic analysis at the molecular level.
Recombinant protein overproduction had a profound and game-changing impact on protein science, making previously inaccessible targets readily available. A very large number of proteins, their mutants and variants have been produced recombinantly, and their structure and function determined at high resolution. The availability of entire genomes has made it possible to address the protein repertoire of cells on a systems-wide scale, applying high-throughput technologies [35]. Recombinant protein expression in *E.coli* as a prokaryotic expression host has become commonplace in molecular biology laboratories world-wide. The recombinant production of protein complexes of eukaryotic origin, however, poses a number of challenges which frequently rule out prokaryotic expression hosts. Eukaryotic proteins are often large and can exceed the size range *E.coli* can overproduce efficiently (typically up to ~100kDa). Post-translational modifications and processing are commonplace in eukaryotic proteins and can be essential for activity, but are generally not supported by a prokaryotic host. The eukaryotic protein folding machinery differs significantly from the chaperone system in *E.coli*, further restricting its utility for eukaryotic protein production. Much effort has been and is being devoted to improving prokaryotic host systems for heterologous production [36-38]. However, in many cases eukaryotic proteins and their complexes will likely require a eukaryotic expression host system for their overproduction, and if a eukaryotic system can be applied with comparable ease as *E.coli* based expression, then this system will likely be a preferred choice. The MultiBac system has been developed precisely with this intention to put in place such a eukaryotic expression system that supports high-level and high-quality production of eukaryotic proteins and their complexes, by using standard operating protocols (SOPs) which make its application comparably facile and routine as *E.coli*-based expression [18, 39-41].
2.1/ MultiBac developments

The baculovirus / insect cell expression system is particularly well-suited for the production of eukaryotic proteins. At the core of this expression technology is a recombinant baculovirus into which the heterologous genes of interest have been inserted. This composite baculovirus is then used to infect insect cell cultures grown in the laboratory. MultiBac is a more recent baculovirus / insect cell system which has been specifically tailored for the overproduction of eukaryotic complexes that contain many subunits [40]. An important prerequisite for the efficient expression of eukaryotic proteins and their complexes are easy-to-use reagents for (multi)gene assembly and delivery. Equally required are robust and standardized protocols for all steps involved in the expression experiment, from gene to purified protein complex. These steps should ideally be implemented as standard operating procedures (SOPs), especially in laboratories where the expression experiment itself and its optimizations are not the primary objective, but rather the protein complex and the determination of its structure and mechanism within a reasonable time-frame. The implementation of such SOPs will then enable non-specialist users to apply the technology with relative ease. The MultiBac system has been designed to meet these requirements [18, 40, 41].

MultiBac consists of an engineered baculovirus that has been optimized for multiprotein complex expression [17] (Fig. 1). The MultiBac baculovirus exists as a bacterial artificial chromosome (BAC) in *E.coli* cells (DH10MultiBac or DH10MB). The replicon (F-factor) present on the BAC restricts its copy number to (typically) one [42]. The MultiBac genome has been modified by deleting proteolytic and apoptotic functionalities from the baculoviral genome that were found to be detrimental for the quality of the heterologous target complexes produced [17-19, 23, 41]. The MultiBac system furthermore comprises an array of small custom-designed DNA plasmid modules that facilitate the assembly of multigene expression cassettes and their integration into the baculoviral genome (Fig. 1).
Integration of the multigene expression cassette constructions into the baculoviral genome occurs via two sites (Fig. 1). One is a mini-Tn7 attachment site embedded in a lacZα gene that is used for blue/white selection and is accessed by the Tn7 transposase which is expressed in the DH10MB cells from a helper plasmid as described previously [43]. Upon integration into this Tn7 site, the LacZα gene is disrupted; white colonies indicate successful transposition. A second entry site is formed by a short imperfect inverted repeat, LoxP, at a location distal from the Tn7 attachment site (Fig. 1). It can be accessed by means of the Cre enzyme, a site-specific recombinase that targets the LoxP imperfect inverted repeat (Fig. 1). Cre integration occurs by fusing LoxP sites present on the MultiBac genome on the one hand, and on a DNA plasmid module on the other. Successful Tn7 and also Cre integration is imposed by antibiotic selection against the resistance markers encoded by the DNA plasmid modules integrated into the MultiBac genome. The integration sites can be used to integrate genes encoding for one or several multiprotein complexes of choice, but also for additional functionalities that may be required to activate or inactivate the complex (kinases, phosphatases, acetylases, deacetylases, others), support its folding (chaperones) or post-translational processing such as glycosylation [19, 23, 44-46].

The composite MultiBac baculoviral genome which contains all desired heterologous genes is then purified from small bacterial cultures using standard alkaline lysis protocols and applied to small insect cell cultures, typically in six-well plates, with a lipidic or non-lipidic transfection reagent [24, 41]. The resulting live baculovirions are harvested and applied to larger insect cell cultures for heterologous protein production and purification. Production baculovirus is then stored for example at 4°C in the dark to avoid degeneration of viral titers. A more secure long term storage method is provided by freezing small aliquots of baculovirus infected insect cells (BIICs) that are then stored in liquid nitrogen [47].
A centerpiece of the MultiBac system is the facilitated assembly of genes into multiexpression cassettes (Fig. 1, 2). Originally, this was addressed by creating two different DNA plasmid modules that contained a so-called Multiplication Module. This module allowed iterative assembly of single or dual expression cassettes, each fitted with a promoter (p10 or polh) by restriction/ligation utilizing compatible sites that would be destroyed upon ligation [17]. One plasmid (pFBDM) accessed the Tn7 site, the second plasmid (pUCDM) accessed the LoxP site on the MultiBac genome. Both plasmids could be fitted with one to many foreign genes by means of multiplication. The pUCDM plasmid contains a conditional origin of replication derived from the phage R6Kγ. Its survival therefore hinges on the presence of a particular gene (pir) in the genome of specific E.coli strains. If a pUCDM plasmid is transformed into DH10MB cells which are pir negative, it will only survive if it is productively fused to the MultiBac genome by Cre-LoxP reaction [17].

The plasmid module repertoire of the MultiBac system was subsequently expanded by fitting out the pFBDM plasmid with a LoxP site, giving rise to pFL. A further plasmid was created, pKL, which in contrast to the high-copy number pFL plasmid is propagated at low copy numbers, and thus could accommodate difficult or very large genes that had turned out to be unstable in pFL. A new version of pUCDM was designed with a different resistance marker (spectinomycin), pSPL. pSPL or pUCDM could now be fused either with the MultiBac genome in DH10MB if Cre was expressed, or in vitro with pFL or pKL. All plasmids retained the Multiplication Module and therefore could be outfitted with several to many genes in an iterative way. pUCDM and pSPL were now denoted Donor plasmids (D), while pFL and pKL were denoted Acceptors (A) (Fig. 2) [18, 41]. One Acceptor could be fused with one or two Donors by Cre-LoxP reaction in vitro, and productive fusions were identified by the combination of resistance markers present on the fusion. The fused AD or ADD constructs carrying several to many heterologous genes, could then be inserted into the
Tn7 attachment site of the MultiBac genome by transposition in DH10MB cells as before. *In vitro* fusion of AD or ADD plasmids prior to integration into the Multibac genome did not rule out use of the LoxP site present on the viral backbone. Integration into the viral loxP site simply had to precede the transposition reaction [18].

A *sine qua non* of contemporary structural biology is automation, to increase the throughput of expression experiments by robotic approaches. As a consequence, the multigene assembly technology of the MultiBac system was adapted to automation by using a liquid handling robot [24, 25]. For this, the tandem recombineering technology (TR) originally developed for prokaryotic complex expression [38] was adapted to the MultiBac system [48]. TR combines sequence-and-ligation independent gene insertion (SLIC) with Cre-LoxP fusion to generate multigene expression constructs in high-throughput in a robotic setup. Adaptation of the MultiBac plasmids to TR required subtle adjustments of the plasmids resulting in Acceptor plasmids pACEBac1 and pACEBac2 as well as Donor plasmids pIDK, pIDS and pIDC that are robotics-compatible [22, 24, 40]. Concomitantly, with the objective to simplify the monitoring of protein production by measuring fluorescence levels, a new baculovirus genome, EMBacY, was created expressing a fluorescent protein (YFP) from its backbone, with fluorescence intensity increasing concomitantly with the quantity of the heterologous protein complex expressed at the same time. All Acceptor and Donor plasmids developed for the MultiBac system over time are compatible with each other, and also with the MultiBac and EMBacY genomes (and other MultiBac genome derivatives) that were and are being developed (see also 2.3 and 2.4).

**2.2/ A MultiBac user access platform**

The MultiBac system rapidly developed into a sought after tool following its introduction and original publication, compellingly underscoring the current need for an accessible
multiprotein expression technology for eukaryotic multiprotein complexes. The number of laboratories using MultiBac approaches a thousand at the time this review is written. Research groups in academia, the biotech and pharma industries are implementing MultiBac to produce their specimens of interest. Moreover, biotech spin-offs were founded based on MultiBac developments, including a preclinical vaccine development company, Redvax GmbH, which in 2015 was acquired by the global pharma enterprise, Pfizer.

The high demand for accessing this technology resulted in the establishment of a dedicated research and training platform at the European Molecular Biology Laboratory (EMBL) in the Eukaryotic Expression Facility (EEF) established at the EMBL Outstation in Grenoble (Fig. 3). The EEF has been supported since 2008 by the European Commission (EC) through infrastructure grants (P-CUBE, BioStruct-X, INSTRUCT) and by the national (French) research agency (ANR) through the Investissement d’Avenir program. More than a hundred projects per year, by local national and transnational users, including academic research projects as well as industrial contracts have been processed in the facility, covering a wide range of applications in basic and applied research and development. Academic project access is based on the sole criteria of excellence, determined by an independent panel reviewing research proposals. The facility has implemented a SOP-based procedure for routinely and rapidly moving projects through the MultiBac platform pipeline [23, 24].

The entire process from preparing the multigene expression construct to (small-scale) purification of the specimens of interest requires around two weeks according to this protocol. A large number of constructs can be processed in parallel according to this protocol. Virus amplification as well as heterologous protein production is monitored by measuring the signal of yellow fluorescent protein (YFP) in small (1 million cells) aliquots withdrawn from the cell cultures at defined intervals. The YFP signal reaching a plateau indicates maximal production of the desired complex specimen and the expression cultured is harvested at this
point for further processing and purification. Particular care is taken to assure highest quality virus production during virus amplification – early (budded) virus is harvested throughout the amplification process to avoid accumulation of defective viral particles that would compromise virus performance. Initial virus is stored at 4°C, while production virus is stored as BIICs in liquid nitrogen [21-24] (Fig. 3).

2.3/ OmniBac: Universal multigene transfer vectors

Two approaches for foreign gene integration into the baculoviral genome dominate the field. One approach depends on the baculoviral genome present as a BAC in *E. coli* cells, and relies on gene integration by transposition catalyzed by the Tn7 transposase which is constitutively expressed from a dedicated (helper) plasmid in the cells harboring the BAC. The foreign genes are provided by transforming a transfer plasmid into these *E. coli* cells. Selection for recombinant virus occurs based on the resistance marker that is present on the transfer plasmid and also integrates into the BAC that is the baculoviral genome, and blue / white selection – productive transposase mediated integration destroys a lacZα gene. The MultiBac system applies this strategy, extended by the option to provide in addition to the Tn7 transposase also the site-specific Cre recombinase from a second dedicated helper plasmid, to fuse an additional plasmid into a LoxP site present on the MultiBac baculoviral genome [17, 40, 41].

The alternative, original approach is based on homologous recombination, mediated by DNA regions in the transfer plasmid that are also present in the baculoviral genome, flanking the polh locus that had been inactivated by destruction of the *polh* gene. These regions of homology correspond to the open reading frames Orf1620 and lef2/603. By using this method, insertion occurs by co-transfecting the transfer plasmid and the purified baculoviral genome into insect cells. The baculoviral genome is typically linearized in the
region between Orf1629 and lef2/603, to suppress formation of virus devoid of the heterologous DNA of interest. Fusion of the plasmid DNA with the baculoviral DNA to create a replicating genome is then achieved via the homologous recombination system of the insect cells. The genome is thus re-circularized, concomitantly inserting the gene of interest. Live virions are then produced that express the desired protein(s).

Both methods have advantages and disadvantages. The Tn7/BAC based integration approach is the method of choice in many laboratories, mainly due to its simplicity. Since the baculoviral genome exists as a BAC in E.coli, it can be, in theory, propagated indefinitely and used for many experiments after obtaining the initial aliquot. Moreover, it can also be manipulated by gene editing technologies in its E.coli host cell. In contrast, the linearized baculoviral DNA for homologous recombination has to be obtained from the supplier for every experiment *de novo* and cannot be propagated at will. Furthermore, the homologous recombination method is arguably somewhat more involved and may require specialist knowledge. On the other hand, this approach is more amenable to automation as compared to the BAC-based method which is characterized by many steps some of which (such as blue/white screening) cannot be readily scripted into robotics routines. A further disadvantage of the BAC-based system may be found in the relative instability that was found for BAC-derived baculoviruses in insect cells, presumably originating from the presence of extended bacterial DNA elements (selection marker, F-replicon, LacZα gene) [4, 49], limiting its use in human applications mainly to preclinical studies [4]. In fact, baculoviruses used in commercial production to date are still being made almost exclusively by applying the classical homologous recombination technique [4].

Available BEVS all relied on either one method or the other, which were mutually exclusive, and the choice of the transfer plasmid decided which virus would be used for protein production. This situation is unsatisfactory given that both systems (BAC/Tn7-based
and classical) provide unique opportunities. Moreover, numerous baculoviruses with customized functionalities have been created for both applications, each with its own merit. We therefore designed and created the “OmniBac” transfer plasmids which combine the DNA elements required for Tn7 transposition in the BAC-based system and also the homology regions for baculovirus generation following the classical method, and thereby are universally applicable (Fig. 4). These OmniBac transfer plasmids function as Acceptors in the MultiBac system (Fig. 4). They can be combined with a variety of Donors to yield multigene Acceptor-Donor fusions that can then be funneled into any baculovirus of choice [50].

2.4/ The ComplexLink polyprotein technology

A challenge frequently encountered when overexpressing protein complexes relates to unbalanced expression levels of the individual protein subunits that are to assemble into the biological target superstructure. If a particular subunit is badly made in a co-expression experiment, it will limit the overall yield of the fully assembled protein complex dramatically. This challenge can be addressed within limits by co-infection approaches using several viruses, or the choice of promoters. In contrast to very late promoters such as polh and p10, earlier promoters are expressed at lower levels. However, it may be often impractical to resort to co-infection or to tuning protein expression levels by promoter choice, also due to the fact that the timing of the production of protein subunits will then be altered as well, with often unpredictable consequences. A solution to such problems can derive from observing the strategies certain viruses utilize to realize their proteome. Coronavirus, the agent that causes severe acute respiratory syndrome (SARS), produces its complete proteome from two long open reading frames (ORFs) that give rise to polyproteins in which the individual protein specimens are covalently linked. A highly specific protease, also encoded by the ORF, then liberates the individual proteins by cleaving them apart.
The ComplexLink technology implements this strategy for recombinant polyprotein production from polygenes [22, 40, 51] (Fig. 5). In ComplexLink, genes encoding for a protein complex of choice are conjoined to yield a single ORF. This ORF gives rise to a polyprotein in which the individual proteins are linked by short amino acid sequences representing a cleavage site for the Nia protease from tobacco etch virus (TEV) which is also encoded by the ORF and is the first protein produced. In addition, a cyan fluorescent marker protein (CFP) is present at the C-terminal end of the polyprotein construction. Upon translation, TEV protease liberates itself, and all other proteins including CFP by cleaving the TEV-specific proteolytic sites. The ComplexLink plasmids, pPBac, pKL-PBac and pOmni-PBac function as Acceptors in the MultiBac system and can be fused to Donors which may contain further genes encoding for polyproteins. The ComplexLink technology proved to be highly successful in producing difficult-to-express protein complexes in high quality and quantity, including the physiological core complex of human general transcription factor TFIID [52-54]. A notable exploit is influenza polymerase, an important drug target to combat the flu, which has remained inaccessible for 40 years since its discovery. Influenza polymerase has been produced, for the first time, successfully using ComplexLink in conjunction with MultiBac, enabling elucidation of its structure and mechanism by X-ray crystallography at near-atomic resolution [55, 56] (Figs. 5, 6).
3/ Applications
The MultiBac system in its original form was introduced in 2004, and has become the method of choice for a wide range of applications. Primarily developed for accelerating structural biology of multiprotein complexes, it has since then been modified and improved to benefit also other fields, in basic and applied research. We have followed these developments with interest and have occasionally highlighted them in invited review articles and commentaries [40, 53, 57, 58]. In the following, we intend to summarize these developments without being exhaustive, focusing on recent exploits by researchers who adopted the system we had developed, to catalyze their research.

3.1/ Accelerating complex structural biology
The first community which implemented MultiBac were structural biologists interested in elucidating the architecture and mechanics of important multiprotein machines in cell biology at the molecular, near-atomic level. This is also the field were an impressive flurry of highest impact MultiBac-enabled contributions was achieved to date. We had highlighted some of these exciting structures, including important drug targets such as the LKB1-STRAD-MO25 complex [59], and the first structure of a nucleosome-bound chromatin remodeler, Isw1 [60] in a contribution just over two years ago in Trends in Biochemical Sciences [40]. In the short time since then, spanning a mere two years, a large number of new structural studies have been carried out using material produced with MultiBac. A selection of these exploits is presented in Figure 6.

Landmark achievements are the recent crystal structures of influenza polymerase [55, 56]. This success was catalyzed by applying the ComplexLink and MultiBac technologies together, to produce this trimeric protein complex which had remained elusive for decades. The structures describe fluA and fluB variants of the polymerase bound to its RNA ligand, and provide important structural insights into cap-snatching and RNA synthesis by this
enzyme complex, opening up new avenues for pharmaceutical development to combat flu.

Further crystallographic exploits include the structures of human cytoplasmic dynein-2 primed for its power stroke [61], the human argonaute-2 / RNA complex [62], the structure of the spliceosomal protein Prp8 bound to and RNA helicase, Brr2 [63, 64], structures of PI4KIIIβ kinase complexes [65] among numerous others. Highlights achieved by using MultiBac produced material for electron microscopic studies include the structures of COP1-coated vesicles, revealing alternate coatomer conformations and interactions [66,67], the architecture of the physiological core of human general transcription factor TFIID [52, 58], or the elucidation of the molecular mechanisms of the anaphase promoting complex APC/C at sub-nanometer resolution [68]. Recently, the entire human Mediator transcription factor holocomplex has been successfully assembled by using MultiBac, and functionally characterized [69]. MultiBac reagents have been incorporated into pipelines for producing membrane proteins and their complexes [70]. We anticipate that many more exciting structures of important protein assemblies will be determined in the future, by using the MultiBac system as a production tool.

3.2/ MultiBac in pharma and biotech

The baculovirus / insect cell system has had a major impact on the production of high-value pharmacological targets, for pharmaceutical characterization, structure-based drug design, diagnostics, biosensor engineering and high-throughput proteomics [2, 4, 5, 71]. Notably human proteins, virus-like particles and vaccines have been successfully expressed by BEVS [2, 4, 5]. Glycoproteins are sought-after biologics in the phama and biotech sector, and insect cells have proven to be well suited for the expression of biologically active and immunogenic specimens. The MultiBac system has been engineered to enable high-quality production of glycoproteins and their complexes. The original MultiBac baculoviral genome was already lacking the $v$-cath and chiA genes, which are encoding for a cathepsin-like protease and
chitinase, respectively. Both v-cath and chiA have been shown to be detrimental to glycoprotein production [72-74]. The glycosylation pattern of secreted proteins in insect cells differs from mammalian patterns which involve more complex N-glycans [75, 76]. These differences can have detrimental effects on human glycoproteins produced in insect cells. To overcome this impediment, a new MultiBac-derived baculovirus, SweetBac, was constructed, which includes glycosylases in the backbone, resulting in mammalianized glycosylation patterns of SweetBac produced glycoproteins [23, 44, 45]. More recently, improved MultiBac variants were introduced to minimize fucosylation in insect cell derived glycoproteins to reduce binding to antibodies from the sera of patients with allergies [46].

The baculovirus / insect cell system has demonstrated its aptitude to produce complex multicomponent assemblies such as virus-like particles (VLPs) [4, 9-11, 77]. VLPs are promising candidates for vaccination. VLPs resemble natural virus shells, but are lacking genetic material and therefore are safe and not infectious. VLPs can be proteinaceous, such as for example papilloma VLPs used to prevent cervical cancer. More recently, enveloped viruses have been successfully produced using BEVS, including influenza and chikungunya vaccine candidates [4, 77-80]. The MultiBac system was already successfully used to produce a number of VLPs including an array of papilloma serotypes [40]. Complex VLPs representing highly pathological virus strains were produced safely by MultiBac [81]. In particular the availability of OmniBac plasmids, which are part of the MultiBac vector suite, may provide unique opportunities for VLP vaccine development, as they are and are equally useful for exploratory preclinical studies in high-throughput as well as pharmacological production by choosing the most appropriate viral backbone for large scale expression.

Baculoviruses not only infect insect cells, but can also transduce mammalian cells efficiently [15, 82-84]. By choosing mammalian-active promoters instead of polh, p10 or other baculoviral promoters, proteins of interest can be produced from a baculovirus that has
entered a mammalian host cell. Baculoviruses do not replicate in mammalian cells, therefore, the current consensus is that this BacMam approach can be performed safely in laboratories. A particular benefit BacMam is that large DNA insertions including multicomponent signaling cascades or entire metabolic pathways can be transduced into mammalian cells efficiently by the baculovirus which can tolerate very large gene insertions, and can be amplified and produced in large amount in a straight-forward way in insect cell cultures. Baculovirus is thus emerging as a highly promising gene delivery tool into mammalian cells, for a multitude of applications [14, 15]. In this context, multigene MultiBac constructions were successfully used to produce recombinant adeno-associated viruses (AAVs) for gene therapy [12, 13, 40].

3.3/ Synthetic Biology: Rewiring the genome

The AcMNPV genome has a size of around 130 kilobases and contains many functionalities which are essential in the natural life cycle of the virus, but dispensable in laboratory culture. Moreover, in the laboratory, it has been recognized that the genome has a tendency to undergo multiple deletions in its genome during amplifications, notably in regions containing foreign gene cargo that has been inserted for overproduction [4, 50]. This can have severely detrimental consequences for heterologous target protein production, especially for fermenter-scale expression of biologics which require large foreign DNA insertions in the viral genome, and several rounds of amplification until sufficient volumes of production virus are obtained to charge the fermenter. We have shown that some of these limitations can be overcome at least on laboratory-scale by stringently adhering to virus generation protocols that avoid or limit the occurrence of widespread deletions [18]. Moreover, it appears that there is considerable scope for improving virus performance by eliminating mutational or deletion hot-spots in the viral genome, and by removing DNA regions which are not required in cell culture. The baculoviral genome, in particular when present as a BAC, lends itself
excellently to genome manipulation and editing techniques. We and other have exploited this to remove unnecessary or undesired functionalities from the virus, such as the *polh*, *p10*, *v-cath* and *chiA* genes.

The advent of efficient synthetic biology techniques now holds the promise to reverse this somewhat cumbersome top-down approach, and to rationally redesign and rewire the baculoviral genome bottom-up by applying DNA synthesis and assembly methods that have become available more recently. Interestingly, comparison of baculovirus sequences in genome databases indicated that most genes and DNA elements thought to be required for survival of the virus in cell culture are confined to roughly one half of the circular viral genome, while the other half contains DNAs that can be probably disposed of in the laboratory [50]. This approach may hold challenges given the complex interplay of baculoviral proteins and their relative expression levels during the different phases of the viral life cycle [4]. Notwithstanding, synthetic approaches, probably best applied in combination with sequential deletions, are exciting and avenues for developing new and minimal baculoviral genomes that can be customized for optimal properties the laboratory and also in industrial manufacturing.
4/ Outlook

Since the pioneering first reports more than three decades ago, the baculovirus / insect cell expression system has developed into a mainstream production platform accelerating a wide range of research projects in academic and industrial laboratories. In the post-genomic era, multiprotein complexes have entered center stage as essential catalysts of cellular activity, and notably the MultiBac BEVS has contributed substantially to make hitherto inaccessible protein complexes available, to unlock their structure and mechanism at molecular detail. Moreover, BEVS has emerged as remarkably useful tool in the biotech and pharma sector, for the production of complex biologics in disease prevention and therapy. The development of this versatile expression tool is continuing unabated as it is set to benefit markedly from powerful new synthetic biology techniques that are becoming available. Fueled by these innovations, BEVS is excellently positioned to play a key and increasing role in the life sciences, in basic and applied research in the future. Exciting times are ahead of us.
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Figures Legends

Figure 1. The MultiBac baculovirus / insect cell expression system. The MultiBac system is shown in a schematic view (left). MultiBac consist of an engineered baculovirus optimized for protein complex production. The MultiBac baculoviral genome exists as a bacterial artificial chromosome (BAC) in *E.coli* cells. It contains two integration sites for foreign genes, by transposition or, alternatively, by site-specific recombination mediated by the Cre enzyme. MultiBac further consists of an array of plasmids called Acceptors and Donors. MultiBac baculovirions (centre) are generated by transfection of the composite MultiBac BAC in insect cells. MultiBac is used for a wide variety of applications in basic and applied research and development (right).

Figure 2. MultiBac Acceptor and Donor plasmids. A variety of entry plasmids to integrate heterologous genes into the MultiBac baculoviral genome have been created since the introduction of the system 2004, each with its own merits. Functional modules contained in the plasmids are denoted (bottom). All plasmids are compatible with each other and can be used in various combinations to generate recombinant MultiBac baculoviral genomes for multiprotein expression and/or multigene delivery.

Figure 3: MultiBac expression platform at the EMBL Grenoble. The standard operating protocol (SOP) utilized to express proteins and their complexes by using MultiBac is illustrated. The entire process takes two weeks from generation of the composite MultiBac BAC (bacmid) to quantitative expression analysis. A MultiBac baculovirus variant called EMBacY is used, producing yellow fluorescent protein (YFP) to track virus performance and heterologous protein production. Protein production is monitored by Western blot (WB) or by gele electrophoresis (SDS-PAGE). Production virus is stored long-term in frozen aliquots of baculovirus in insect cells (BIICs).
Figure 4. OmniBac – Universal transfer plasmids for every BEVS. (a) Acceptor plasmids pOmniBac1 and pOmniBac2 are shown schematically, elements are marked above (Fig. 2). These Acceptors can be combined with the MultiBac Donor plasmids by Cre-LoxP recombination. (b) OmniBac plasmids contain elements for homologous recombination and also Tn7 based transposition. Multigene constructions based on OmniBac plasmids can therefore access all available baculovirus genomes. Thus, with the same plasmids, composite baculovirus for preclinical studies as well as manufacturing can be produced efficiently.

Figure 5. ComplexLink technology. (a) The ComplexLink technology was created to produce multiprotein complexes from self-processing polyprotein constructs as shown here schematically. The polyprotein contains a TEV protease and a fluorescent protein at the N-terminus and C-terminus, respectively. Polyproteins are processed during expression into the individual protein entities by the highly specific TEV protease. (b) Polyprotein expression plasmids pPBac, pKL-PBac and pOmni-PBac are shown. DNA modules are marked as above (c.f. Fig 2). (c) Schematic representation of a self-processing polyprotein before TEV-mediated cleavage encoding for influenza polymerase. TEV stands for tobacco etch NLa protease, PA, PB1 and PB2 are subunits of the trimeric enzyme complex, CFP stands for cyan fluorescent protein. BstEII and RsrII are asymmetric restriction enzymes that can be used to cleave polyprotein expression plasmids for restriction / ligation-based heterologous gene integration.

Figure 6. MultiBac complex structure gallery. Selected examples of recent high impact structures of MultiBac produced biological specimens are shown. Examples include cryo-EM architectures of COPI-Coated Vesicles (EMD-2084 to EMD-2088), the complete human APC/C complex with coactivator and substrate (EMD-2651 to EMD-2654) and the human core-TFIID complex (EMD-2229 to EMD-2231). Notable structures that were determined by X-ray crystallography (PDB identifiers are provided in in brackets) include influenza
polymerases A and B bound to the viral RNA promoter (PDB identifier 4WSA, 4WRT), human Argonaute Ago2 in complex with miR-20a RNA (4F3T), the spliceosomal complex Brr2HR / Prp8abl (4KIT), human GABA(B) receptor (PDB 4MQE), the dynein-2 motor bound to ADP (4RH7), the mitotic checkpoint complex MCC (4AEZ) and the GluN1/GluN2B N-methyl-D-aspartate receptor (3QEL). Molecular illustrations were prepared with PyMOL (www.pymol.org) and Adobe Photoshop Version CS6.
Figures

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