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## Objective

The objective is to create a technology landscape report in the area of "Production of Therapeutic Glycoproteins"

### Comment

The idea is to find out the innovative market players in this technology area, those are the players that are the most technologically advanced in the market place. Intellectual Property that these players hold is taken as a measure of their technological prowess in this particular area. We have used Patents to point out the innovative and technologically advanced market players.

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# Methodology



## Introduction

### **Glycosylation:**

Post-translational modifications ? and in particular glycosylation ? play a decisive role in ensuring proper protein function. Glycosylation has been found to occur in a large variety of protein classes, including ? but not limited to ? antibodies, protein hormones, growth factors, cytokines and vaccines. Glycosylation is the covalent attachment of a carbohydrate component to the polypeptide backbone. For some proteins glycosylation can increase solubility, influence biological half-life and/or biological activity. The two main classes of glycosidic bonds to polypeptide chains are termed N-linked and O-linked.

- N-linked glycosylation: the glycan is linked to nitrogen in the side chain of asparagine. The oligosaccharide is attached via the amide
- nitrogen of an asparagine residue in the consensus sequence "asparagine ? any amino acid except proline ? serine/ threonine'
- O-linked glycosylation: the glycan is linked to oxygen in the side chain of mostly serine or threonine, as well as hydroxylysine. No specific consensus sequence has been identified for O-linked glycosylation.

In the case of humans, the oligosaccharides are typically composed of the following monosaccharides: mannose (Man), glucose (Glc), N-acetylglucosamine (GlcNAc), fucose (Fuc), galactose (Gal), N acetylgalactosamine (GalNAc), and sialic or neuraminic acids. The monosaccharides are linked together in a linear or branching fashion via either ?- or ?-glycosidic bonds between specific carbon atoms of the individual monosaccharides.Walsh G et al, Nature Biotechnology,2006

#### Importance of Glycosylation in Therapeutic Proteins:

Many important therapeutic proteins are glycosylated including epoetin (EPO), granulocyte macrophage colony stimulating factor (GM-CSF) and tissue plasminogen activator (t-PA). Following table lists glycosylated therapeutics currently on the market.

### Table I. Glycosylated therapeutics approved by the FDA and/or EMEA

Product	Class	Mode of action	Indication	Cel
Aranesp (Darbepoetin alfa)	Erythropoiesis stimulating protein	Regulates red blood cell production	Anemia	CH
Arcalyst (Rilonacept)	IL-1 Trap	Binds IL-1β to prevent the interaction to cell surface receptors	Cryoporin-associated periodic syndromes	CH
Avastin (Bevacizumab)	rMab	Binds to the vascular endothelial growth factor (VEGF) to inhibit angiogenesis	Colorectal cancer	CH
Avonex (Interferon β-1a)	Interferon	Binds to type I interferon receptors to activate two Jak tyrosine kinases	Multiple sclerosis (MS)	CH
Cerezyme (Imiglucerase)	Enzyme	Engineered to have mannose-terminated oligosaccharide chains that are recognized by endocytic carbohydrate receptors on macrophages. Catalyzes hydrolysis of glycolipid glucocerebroside to glucose and ceramide in those macrophages which accumulate lipids in Gaucher disease	Gaucher Disease	CH
Elaprase (Iduronate sulfatase)	Enzyme	Hydrolyzes the 2-sulfate esters of terminal iduronate sulfate residues from the glycosaminoglycans dermatan sulfate and heparan sulfate in the lysosomes of various cell types	Hunter syndrome	НТ
Enbrel (Etanercept)	Fusion protein	Mimics inhibitory effects of naturally occurring soluble TNF receptors to reduce inflammatory response	Rheumatoid arthritis	CH
Epogen (Epoetin alfa)	Erythropoiesis stimulating protein	Recombinant human erythropoietin interacts with erythropoietin (EPO) receptors to stimulate production of red blood cells from bone marrow stem cells	Anemia	CH
Erbitux (Cetuximab)	rMab	Binds to the extracellular domain of the epidermal growth factor (EGFR) preventing activation of EGFR to impair cell growth and proliferation	Colorectal cancer	SP
Herceptin (Trastuzumab)	rMab	Binds to HER2+ tumor cells, blocks downstream HER2 signaling to inhibit proliferation of cells	Breast cancer	CH
Wellferon (Interferon α)	Cytokine	Binds to type I interferon receptors (IFNAR1 and IFNAR2c) which, upon dimerization, activate two Jak (Janus kinase) tyrosine kinases (Jak1 and Tyk2). Upregulates expression of MHC I proteins	Chronic hepatitis C	Hu
Mircera (Methoxy polyethylene glycol-epoetin	Erythropoiesis stimulating protein	Recombinant human erythropoietin interacts with erythropoietin (EPO) receptors to stimulate production of red blood cells from bone marrow	Anemia	CH

List of Glycosylated Therapeutic Proteins Hossler P et al, Glycobiology,2009 [1] Glycosylation is important because it can influence the biological activity of a protein through different mechanisms. First of all, glycosylation can affect half-life by influencing the active clearance of a protein. Certain pituitary glycoprotein hormones such as lutropin are rapidly removed from circulation by the liver if they bear a sulphated GalNAc residue. A second example is EPO, the serum half-life of which is dependent on four sialylated N-glycans (sialic acid is itself a glycan). Poorly glycosylated forms of EPO are rapidly cleared by filtration in the kidney. In addition, under-sialylated EPO is rapidly removed by hepatocytes and macrophages. Thus, increasing the degree of sialylation and glycosylation decreases the renal clearance rate, and can increase EPO in vivo activity.

Glycosylation can affect the activity of a recombinant protein more directly. Rituximab, for instance, is a monoclonal antibody (mAb) used to treat non-Hodgkin?s lymphoma through targeting of B-cell CD20 antigens. Its activity can be assessed with the antibody-dependent cellular cytotoxicity (ADCC) test, which measures the ability of an engineered mAb to lyse target cells. The ADCC of two types of mAb (one of which was rituximab) was





A protein with typical post-translational modifications and functional consequences Kullmann M et al, Nephrol Dial Transplant. 2006[2]



Schematic view of control methods in the glycosylation pathway in a CHO cell line Hossler P et al, Glycobiology, 2009[3]

#### **Glycosylation and Biopharmaceutical Manufacturing:**

Protein glycosylation typically leads to a diverse array of product glycans due to multiple factors. Even with knowledge of human glycoform profiles for a particular glycoprotein, matching those attributes on the recombinant biologic is not trivial. Regulatory agencies around the world require biopharmaceutical companies to both characterize and maintain product quality attributes (including glycosylation) within defined acceptance limits. In biopharmaceutical manufacturing, the acceptance limits are based on what the cell line, upstream cell culture, harvest operation and downstream purification can support, and on the known glycoform species.

Current production processes usually use microbial systems, yeast or cell lines derived from insects (SF9), mice (SP20) or hamster (CHO). Glycosylation is notably sensitive to cell growth conditions. Changes in culture pH, the availability of precursors and nutrients, and the presence or absence of various cytokines and hormones can each affect the extent of glycosylation. In addition, the presence of sialidases and other glycosidases, either secreted or released by dead cells, can cause degradation of a previously intact product.

Control of this metabolic pathway can be manifested through a variety of different conditions, including the cell lines, bioprocess, and cell culture media used for protein production. Hossler P et al, Glycobiology, 2009



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Glyco-optimised biotherapeutics Baumeister H et al, Glycotope[4]

#### **Glycosylation in Biosimilars:**

A change is occurring in the biotechnologically derived drug industry with the impending arrival of biosimilars on the market. This is a challenge for biopharmaceutical companies where a cell line change, process changes and/or change of manufacturing site requires extensive comparability studies to prove that the molecule remains the same. Glycosylation has moved centre-stage in the development of biopharmaceutical drug products. Its influence on a protein?s characteristics has been acknowledged by regulatory requirements for the development of biosimilars that demand a high level of glycosylation-based similarity to the original product. As a consequence, glycosylated biosimilars need to be equipped with a similar pattern of glycosylation.Baumeister H et al, Glycotope

## **Relevant Patents' Example**

1	<u>US6780607B2</u>	8/24/2004	Method for cell-free protein complete post-translational modification	The present invention relates to methods of production of the completely post-translationally modified protein by combination of cell-free protein synthesis and cell-free co-and post-translational modification. Previous cell-free protein synthesis system has only been capable of producing partially post-translationally modified protein but the present invention employs a co-and post-translational modification machinery that produces completely post-translationally modified protein.	DreamBiogen Co., Ltd.	For a mammalian protein to be active, it must show native post translational modifications. One way to produce proteins with native post translational modification is to culture the cells in native cell lines or modified eukaryotic hosts. This process is time consuming. Also, the available cell-free protein synthesis systems till then could not produce proteins with complete modifications.	An advanced cell free protein synthesis system is described which works by the combination of cell free protein synthesis and cell free complete post translational modification. The system comprises ribonucleotide triphosphates, sufficient amount of co- and post-translational modification machinery such as ER/golgiaparatus or ER/Golgi apparatus/plasma membrane, or other organelles and DNA or RNA template.
2	<u>US6610516B1</u>	8/26/2003	Cell culture process	A glycoprotein is produced by a process comprising culturing mammalian host cells expressing nucleic acid encoding a glycoprotein in the presence of (a) a factor that modifies growth state in a cell culture, (b) a divalent metal cation that can adopt and prefers an octahedral coordination geometry, and/or (c) a plasma component. In this process, the occupancy of an N-linked glycosylation site occupied only in a fraction of a glycoprotein is enhanced. Such culturing is preferably carried out at a temperature of between about 30° C. and 35° C. and/or in the presence of up to about 2 mM of a butyrate salt and/or in the presence of a cell-cycle inhibitor.	Genentech, Inc.	There is a need for increasing glycosylation site occupancy in glycoproteins having multiple glycoforms to produce glycoprotein therapeutics of consistent product quality.	A few modifications to the cell culture conditions have been described to increase the occupancy of a glycosylation site in productions of glycoprotein with multiple glycoforms. The modifications include addition of butyrate salt or divalent cations such as iron and manganese. Also, the cell culture temperature has been reduced to 30 - 35 degrees C instead of traditional 37 degrees C. These modifications help control the cells in Go/G1 phase of cell cycle. Glycosylated Human t-PA has been produced.

Click here to download analyzed samples

# Taxonomy

```
.markmap-node {
    cursor: pointer;
}
.markmap-node-circle {
    fill: #fff;
    stroke-width: 1.5px;
}
.markmap-node-text {
    fill: #000;
    font: 10px sans-serif;
}
.markmap-link {
    fill: none;
}
```

```
pre, .mw-code{
    background-color: transparent;
}
d3.xml("https://www.dolcera.com/wiki/images/Manufacture_of_Glycosylated_Therapeutic_Proteins2.mm", function(error, data) {
    if (error) throw error;
        markmap("svg#mindmap_3f9a5b3dfc9aa7c92796a8b2alc409ff", data, {
            preset: "colorful",
            linkShape: "diagonal"
        }, "xml");
    });
```

Note: Taxonomy will be further populated as per detailed analysis

# Assignee Analysis and IP Activity

### **Top Assignees**

• Roche Holding Ltd, Kirin Holdings Company Limited and Novo Nordisk are the top assignees in the field.



**Top Assignees** 

### **Geographical Distribution**

Based on Patent No.



### Based on relevant patents

Note: Countries with three or lesser patents have been grouped into Others.

• Including all family members



### Based on patent family members

Note: Countries with 15 or lesser patents have been grouped into Others.

### **IP** activity

• Patent activity based on Publication Year



IP activity

· Patent activity based on Priority Year



IP activity

### **Top Cited patents**

• Patents with the maximum number of forward citations were determined and the graph shows the top 10 patents with corresponding assignees.



#### Top cited patents

## Dashboard

Patents were categorized on the basis of patent focus- whether novel or modified cell lines or hosts, transgenic animals, novel enzymes, chemical/cell free methods of synthesis, and novel fermentation methods or culture conditions. Only those patents with method of manufacture of therapeutic glycoproteins as their focus have been considered in this analysis. Click here to access dashboard

D Search in: Title, Abstract, Claims	Patents		d. Charts	
Manufacture of Glycosylated the	Publication	Title		
CHO (86)	US20020045207A1	Glycoprotein production process		
C Other mammalian cell lines,	W02011035884A1 1 Process for producing molecules containing specialized glycan structures			
Caransgenic animals (33)	W0199805550149 Improved method for site-selective dycosylation			
Insect (32)	MY1937008			
Plants (85)	C730000000242			
Cast (114)	C220000803A3 Transgenic mammals producing oligosacchandes in their milk			
C Other Fungi (37)	MX1984568	No bible available		
Bacteria (22)	SK200000304A3	No title available		
Other microbe (6)	MX2000002266A	No title available		
Enzymes (170)				
Chemical synthesis/ Cell fre Fermentation/ Cell culture o Add Edit Delete Get Info Add Edit Delete Get Info A Science (1) GENCY FOR SCIENCE TECHNOOGY # GPLIED PHYTOLOGICS INC (1) RTES Biotechnology GmbH (3) Abbott Laboratories (7) Academia Sinica (1) Advantest Corp (2)	GLYCOPROTEIN PRODUCTIO Priority Date (y-m-d): 1997- First Inventor: KRUMMEN LYNN US Class (primary): 4350691 IPC Class (primary): C12P021 Abstract: Glycoprotein production metho Dolcera Summary Not available	IN PROCESS 10-31 IE A US 1 106 ds and host cell lines are provided. The production methods and	<ul> <li>Chains:         <ol> <li>A process for producing a glycoprotein in a editory omprising the steps of: introducing into the editory capable of being expressed by the eukaryotic of encoding a galactosyltransferase and the second encoding a sialyltransferase; maintaining the editory expression of the galactosyltransferase, the siated of the process according to claim 1 wherein the signature of the process according to claim 2 wherein the second of the proces according to claim 2 w</li></ol></li></ul>	
Idvantest Corp (2)	Not available			
gency Of Ind Sci & Technology (1)				
No Date Filter •				
All Patent Types 🔹	Rating: Tags:	Notes:		
All Taos	Houng:	HULLSI		

Dashboard on "Production of Therapeutic Glycoproteins"

## Top Assignees on the basis of production methods

Mammalian cell lines



Transgenic animals



Top assignees in transgenic animals category

Plants



Top assignees in plants category

• Microorganisms



Top assignees in microorganisms category







Top assignees in chemical/cell free synthesis category

• Fermentation/Culture conditions



# Patent product mapping example

S.No.	Patent/ Pub No.	Title	Assignee/Applicant	Product	Product image
1	<u>US5459031</u>	Methods for controlling sialic acid derivatives in recombinant glycoproteins	Amgen Inc.	<u>Epogen®</u>	

• Note: The patents focusing on methods to produce glycosylated proteins, methods to alter the glycosylation pattern of the proteins, cell lines for producing glycosylated proteins or altering the glycosylation pattern, enzymes for producing glycosylated proteins or altering the glycosylation pattern were considered on target in the analysis. Patents focusing on Purification of glycoproteins or therapeutic use of a glycoprotein were considered as off target.