Erythropoiesis stimulating agents: approaches to modulate activity

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Abstract: Recombinant human erythropoietin (rHuEPO), such as the approved agents epoetin alfa and epoetin beta, has been used successfully for over 20 years to treat anemia in millions of patients. However, due to the relatively short half-life of the molecule (approximately 8 hours), frequent dosing may be required to achieve required hemoglobin levels. Therefore, a need was identified in some anemic patient populations for erythropoiesis stimulating agents with longer half-lives that required less frequent dosing. This need led to the development of second generation molecules which are modified versions of rHuEPO with improved pharmacokinetic and pharmacodynamic properties such as darbepoetin alfa, a hyperglycosylated analog of rHuEPO, and pegzyrepoetin, a pegylated rHuEPO. Third generation molecules, such as peginesatide, which are peptide mimetics that have no sequence homology to rHuEPO have also recently been developed. The various molecular, pharmacokinetic, and pharmacodynamic properties of these and other erythropoiesis stimulating agents will be discussed in this review.

Keywords: darbepoetin alfa, erythropoiesis, erythropoietin, glycosylation, pharmacokinetics, pharmacology, polyethylene glycols

Introduction
Anemia is a condition whereby the number of red blood cells, the amount of hemoglobin in blood, and/or the volume of packed red blood cells is lower than normal. Anemia frequently occurs in patients with renal failure, cancer, and other conditions, such as HIV infection. The treatments for anemia include transfusion, iron supplementation, and dosing with erythropoiesis stimulating agents (ESAs). Along with the benefits of each of these treatment options, there are potential safety risks that are reviewed elsewhere.1,2 Red blood cell production, the process of erythropoiesis, occurs in the bone marrow and is controlled by the natural hormone erythropoietin (EPO), a 165 amino acid glycoprotein primarily produced by interstitial fibroblasts in the kidney.3 The EPO molecule has a compact globular structure that contains one O-linked and three N-linked carbohydrate side chains that constitute approximately 40% of its mass.3 EPO production is regulated at the transcriptional level by hypoxia inducible transcription factor (HIF) subunits HIF-1α and HIF-2α, part of the cellular oxygen sensory machinery.4 Under normal conditions, the HIF-α transcription factor proteins are produced but are rapidly hydroxylated by HIF prolylhydroxylases and targeted for degradation by the Von Hippel Lindau complex before having the ability to activate transcription of the EPO gene. However, under low oxygen conditions, the HIF-α proteins are not hydroxylated and are therefore stabilized, and able to bind to the DNA regulatory elements (hypoxia regulatory elements) of the EPO gene and
activate EPO gene transcription and consequently protein production. Pharmacologic modulation of the HIF-α proteins has recently been investigated as an approach to treat anemia but will not be discussed in this review.

EPO acts on early erythroid progenitors resident in the bone marrow, along with other cytokines, to promote erythroid progenitor survival, proliferation, and differentiation into mature erythrocytes (Figure 1). Hematopoietic stem cells resident in the bone marrow differentiate into multiple myeloid and lymphoid lineages, including the erythroid lineage. The earliest committed erythroid progenitors are classified as burst forming units-erythroid. It is these erythroid progenitors that upregulate the expression of the erythropoietin receptor (EPOR) and become responsive to EPO. The EPO:EPOR interaction induces signaling cascades that induce the differentiation of these progenitors to form colony forming units-erythroid, which themselves are highly responsive to EPO. EPO induces the expansion and further differentiation of colony forming units-erythroid cells into proerythroblasts and erythroblasts. Erythroblasts extrude their nuclei and form reticulocytes that are released from the bone marrow into the circulation and subsequently terminally differentiate into hemoglobin containing erythrocytes. In healthy humans, erythrocytes have a lifespan of approximately 100 to 120 days.

EPOR is a type-1, single transmembrane receptor that exists in preformed homodimers on the cell surface. Two regions of the EPO molecule have been shown to bind EPOR, one with a high affinity (Km 1 nM) and the other a low affinity site (Km 1 µM). A single EPO molecule is proposed to bind to the high affinity EPOR site first and then bind to the other EPOR molecule through the second lower affinity site. The generation of EPOR protein and subsequent trafficking to the cell surface is an inefficient process with only 1%–10% of total cellular EPOR molecules being trafficked to the membrane. A key accessory protein is Janus kinase-2 (JAK2) which binds EPOR in the endoplasmic reticulum, induces correct protein folding, promotes surface expression, and is essential for EPOR signaling. Binding of EPO to EPOR induces a receptor conformational change, which brings two receptor-associated JAK2 molecules into close proximity. Transphosphorylation of JAK2 results in phosphorylation of tyrosine residues located within the cytoplasmic tail of EPOR, which serve as docking sites for signaling adaptor proteins. EPOR stimulation induces the activation of signal transducer and activator of transcription-5 (STAT5), phosphoinositol 3′-kinase (PI3K), the MAP kinase (MAPK), and protein kinase C (PKC) pathways. These signaling pathways promote the survival, differentiation
and proliferation of the erythroid progenitors. A number of molecules have been implicated in the negative regulation of EPOR signaling including, Src homology region 2 domain-containing phosphatase 1 (SHP-1), and suppressor of cytokine signaling proteins SOCS-1 and SOCS-3. Absence of negative regulation of EPOR signaling is associated with familial polycythemia due to cytoplasmic truncations of EPOR that remove SHP-1 and other suppressor binding sites. Though other receptor complexes have been suggested for EPO, these data are controversial and are reviewed elsewhere.

**Glycosylation of recombinant human EPO (rHuEPO)**

In 1985, two independent groups reported the cloning of the human erythropoietin gene. The use of rHuEPO was approved in 1988 and 1989 in Europe and the USA, respectively for the treatment of anemia associated with renal insufficiency, and subsequently approved for anemia associated with myelosuppressive chemotherapy associated with cancer treatment. rHuEPO protein is produced in genetically engineered Chinese Hamster Ovary cells and has a molecular weight of 30.4 kDa and is composed of ~60% amino acids and ~40% carbohydrates. Like naturally occurring EPO, rHuEPO consists of a 165 amino acid single polypeptide chain and contains three N-linked glycosylation sites at asparagine residues (Asn24, Asn38, Asn83) and one O-linked site at serine residue Ser126. rHuEPO produced in African green monkey kidney cell line (COS1) and Chinese Hamster Ovary cells was shown to have a biologic activity equivalent to that of endogenous human EPO in both in vitro and in vivo assays. Until recently, the use of mammalian cell lines was critical for the production of rHuEPO. This was due to the requirement for specific glycosyltransferases and glycosidases for glycans biosynthesis for the correct glycosylation of rHuEPO. The process of glycosylation, unlike the synthesis of amino acids and nucleic acids, is enzymatic rather than defined by a template, and hence the carbohydrates are diverse with respect to both the number and the linkage patterns of the sugar units. Indeed, the carbohydrate composition of rHuEPO N-linked chains is complex and differs in terminal N-acetylleukaminic acid (Neu5Ac) content, O-acetylation of the Neu5Ac residues, N-acetyllactosamine extensions, and degree of branching. Thus, changes in cell lines, growth conditions, or manufacturing processes can affect the final rHuEPO characteristics, including the generation of microheterogeneity in glycosylation. Carbohydrate addition to proteins can influence many aspects of a protein's properties including molecular stability, solubility, immunogenicity, and in vitro and in vivo biological activity. These properties are described below.

**Molecular stability**

A protein therapeutic needs to be stable in order to maintain activity and reduce the potential for immunogenicity. Glycosylation can play an important role in maintaining molecular integrity and can reduce the potential for proteolysis. For example, the carbohydrate on rHuEPO helps maintain molecular integrity and activity. The in vitro biological activities of asialo-rHuEPO and fully deglycosylated rHuEPO were reduced to 35% and 11% of initial activity respectively upon heat treatment, whereas glycosylated rHuEPO lost no activity. Under denaturing conditions, such as guanidine-HCl, heat, and acidic pH, unglycosylated rHuEPO was more readily aggregated and precipitated, whereas glycosylated rHuEPO was able to refold and remain soluble. Furthermore, the peptide component of rHuEPO was protected from oxygen free radical damage if glycosylated.

**Solubility**

The solubility of hydrophobic proteins can be increased by the addition of highly hydrophilic sialic acid containing carbohydrates. For example, unglycosylated *Plasmodium falciparum* merozoite surface protein 1 (MSP-1) was considerably more insoluble than glycosylated MSP-1 when purified from transgenic mouse milk. Unglycosylated MSP-1 was also found to be significantly more immunogenic in monkeys challenged with a lethal infection of *P. falciparum* when used as a vaccine than glycosylated MSP-1. This may have been due to the formation of aggregates and precipitates of the unglycosylated protein that mounted an elevated immune response compared to the glycosylated protein. Alternatively, the carbohydrate may have “shielded” the protein from immune surveillance.

**Immunogenicity**

In rare cases, antibodies to administered rHuEPO are produced in patients that recognize and neutralize endogenously produced EPO, resulting in the development of antibody mediated pure red cell aplasia (PRCA). Outside the United States, an increase in PRCA incidence was observed between 1998 to 2004 and was associated with the administration of a human serum albumin (HSA)-free EPREX1 (recombinant human epoetin alfa) formulation. The potential causes of the increased immunogenicity of rHuEPO to induce PRCA in these formulations have been discussed in the literature.
and included leachates from syringe rubber plungers, use of silicone, micelle formation, and formation of aggregates through mishandling and use of polysorbate 80 in the formulation. However, the true cause for the increase in PRCA remains unknown. Interestingly, it was reported that the immunoreactivity of anti-rHuEPO antibodies in PRCA patient sera was elevated when carbohydrate was removed suggesting that glycosylation can “shield” the underlying protein sequences from the immune system. A single tri-annenary sialylated complex carbohydrate occupies a large volume, approximately 2542 Å. Therefore, the protein surface area subjected to immune surveillance would be reduced in glycosylated proteins. Indeed, carbohydrates reduced polyclonal and monoclonal antibody immunoreactivity to rHuEPO. Similar immune surveillance “shielding” effects of carbohydrate have been reported in other polypeptides such as asparaginase, wasp venom peptide, and P. falciparum MSP-1 protein. Taken together, these studies suggest that glycosylated therapeutic proteins may have reduced immuno-genicity compared to the non-glycosylated counterparts.

In vitro and in vivo biological activity

Given the large size of carbohydrates, the negative charge associated with sialic acid and the potential for steric hindrance and electrostatic repulsion, it would not be surprising if a glycosylated ligand would have a reduced binding affinity for its cognate receptor. Indeed this has been identified to be the case in rHuEPO. Studies examining the in vitro properties of various glycosylated isoforms of rHuEPO found an inverse correlation between glycan number and receptor binding affinity and in vitro biological activity (Figure 2). Furthermore, desialylated rHuEPO and rHuEPO synthesized in bacteria, thus lacking any glycosylation, bound EPOR with higher affinity and had a greater in vitro biological activity than fully glycosylated rHuEPO. In contrast, the in vivo biological activity of rHuEPO is directly related to the number of carbohydrates and sialic acids the molecule contains, to a maximum of 14. This was discovered by examining changes in hematocrit when groups of mice were dosed with equimolar peptide equivalents of rHuEPO, but with different numbers of carbohydrates and sialic acids (9 to 14). A strong correlation was found between the number of sialic acids the rHuEPO molecules contained and the effect on increasing hematocrit. This was identified to be directly related to the reduced clearance and consequently increased serum half-life of molecules containing additional sialic acids. A 3- to 4-fold increase in half-life was observed when a rHuEPO molecule containing 14 sialic acids was compared to a rHuEPO molecule containing the maximum six sialic acids in mice.

Although many studies have been conducted it is still unclear why glycosylation of rHuEPO reduces the clearance of the molecules. There are several potential routes of rHuEPO clearance that have been investigated including clearance by the kidney, hepatic clearance through the asialoglycoprotein receptor, erythroid degradation of rHuEPO after binding to and internalization by EPOR expressing cells in the bone marrow, and degradation and metabolism in the tissue interstitium. Though rHuEPO can be detected in urine, initial data suggesting that the kidney was the primary route of rHuEPO clearance were not supported by subsequent studies examining rHuEPO clearance in patients and animal models that had renal insufficiency. Indeed, only <5% of radiolabeled epoetin beta was detected in the urine from healthy men, supporting the hypothesis that the kidney is not the primary clearance mechanism of rHuEPO. Another mechanism that has been suggested to play a role in the clearance of rHuEPO is the liver. Hepatic asialoglycoprotein receptor (ASGPR) has been reported to bind to ASGPs with subsequent internalization and degradation. Therefore the rate of desialylation of glycosylated rHuEPO was hypothesized to control clearance. However, evidence for the clearance of rHuEPO through the liver may just be relevant for asialo-rHuEPO as this mechanism has not been supported by other studies. For example, the clearance of rHuEPO in normal and liver cirrhosis patients was not different suggesting that liver ASGPR may not play a significant role in rHuEPO clearance. Other studies have also supported the finding that the liver may not play a key role in asialoprotein clearance: the saturation of the ASGPR with asialoorosomucoid did not delay the clearance of sialylated I-orosomucoid and no accumulation of desialylated glycoproteins or lipoproteins were reported in ASGPR knockout mouse circulation, though reduced clearance rates of asialoorosomucoid or asialofetuin were reported.

The bone marrow has been suggested to be a clearance organ of rHuEPO, in part through its effector function. This has been hypothesized to occur though the binding of EPO to the EPOR on erythropoietic progenitor cells and subsequent receptor mediated endocytosis and degradation in the lysosome. The hypothesis was supported by in vitro studies using cell lines that expressed or did not express EPOR: rHuEPO was only cleared at a significant amount from culture media containing EPOR expressing cells. Studies in patients receiving chemotherapy and animal models also appear to support this route of clearance.
to some degree. However, in vivo pharmacokinetic studies do not support receptor mediated endocytosis as a primary route of clearance. The comparison of half-lives of intravenous administration of rHuEPO and an EPO analog with the receptor binding sites mutated (NM385) found that NM385 had a slightly longer half-life in rats or mice but was cleared at a similar rate to rHuEPO. These studies suggest that receptor mediated clearance may play some role in the clearance of rHuEPO from the circulation but is not the primary route.

It is also possible that the tissue interstitium may play a role in the degradation of rHuEPO. Cells involved in the reticuloendothelial scavenging pathways or lymphatic system may play a role in the degradation or metabolism of rHuEPO. Therefore, the presence of glycosylation on the molecule may result in steric and or charge related inhibition
of tissue penetration and reduce the potential for subsequent degradation.

**Glycoengineering and hyperglycosylation analogs of rHuEPO**

Glycoengineering is the process whereby new N-linked glycosylation consensus sequences are introduced into desirable positions in the peptide backbone in order to generate proteins with increased sialic acid containing carbohydrate to increase the serum half-life of the protein and thus the therapeutic activity in vivo. As described above, rHuEPO contains three naturally occurring N-linked glycosylation sites and a single O-linked glycosylation site. The N-linked glycosylation sites have a tetraantenary structure and can contain up to a maximum of four terminal sialic acids whereas the O-linked sites have a bi-antenary structure and contain a maximum of two terminal sialic acids. N-linked glycosylation initiates with the transfer of a 14-oligosaccharide precursor to the asparagine (Asn) located within the Asn-X-Serine (Ser)/Threonine (Thr) consensus motif where X is any amino acid except Proline (Pro). This consensus sequence is known as the glycosylation sequon. The oligosaccharide undergoes maturation via the action of glycosyltransferases that append additional sugar units to the terminal mannose residues. Glycosylation sequons are necessary but not sufficient for N-linked carbohydrate addition to secreted proteins. Alteration of the adjacent sequences can convert non-functional glycosylation sequons to functional sites, indicating sequence context and/or secondary structure influence glycosylation. Secondary structures required for carbohydrate addition within functional glycosylation sequons are β or Asn-X turns. As glycosylation is initiated prior to protein folding, sites normally buried within the molecule can be glycosylated. However, the resultant proteins may have altered protein structures and/or stabilities due to inhibition of correct protein folding as was observed for some rHuEPO analogs.

In contrast, O-linked glycosylation is initiated by the attachment of a single monosaccharide, usually N-acetylgalactosamine, to a Ser or Thr residue. The sugar is subsequently modified by glycosyltransferases to form a mature O-glycan structure. It is unknown why certain amino acid residues in a protein are selected for O-linked glycan addition as there is no clear consensus sequence for O-linked glycosylation which may instead be defined by secondary structure.

The discovery that sialic acid containing carbohydrate was directly proportional to the serum half-life of rHuEPO and in vivo bioactivity led to the hypothesis that additional carbohydrates on the rHuEPO backbone could further enhance pharmacokinetic (PK) and in vivo activity and reduce dosing frequency. Therefore, the approach of introducing additional N-linked glycosylation sites in rHuEPO to generate molecules with extended half-lives and enhanced biological and clinical activity was investigated and is described below.

**Darbepoetin alfa**

Darbepoetin alfa is a hyperglycosylated analogue of rHuEPO, has been approved for the treatment of anemia associated with chronic renal insufficiency and anemia associated with concomitant administered myelosuppressive chemotherapy, and has been used in the clinic for approximately 11 years. Darbepoetin alfa was created based on studies performed to investigate whether the introduction of additional N-linked glycosylation sequons into the rHuEPO peptide sequence could result in molecules with increased glycosylation and sialic acid content and thereby potentially increase the serum half-life and in vivo potency of the molecule. Additional N-linked glycosylation sequons were introduced into multiple sites within the rHuEPO molecule and the combination of sites at Asn and Valine (Val) resulted in the generation of a hyperglycosylated rHuEPO analog with two additional N-linked glycosylation sites, resulting in a total of five N-linked sites. Darbepoetin alfa was found to be efficiently glycosylated, maintained excellent molecular stability, and had a similar structural conformation to rHuEPO. The additional N-linked glycosylation sites increased the molecular weight of the molecule from 30.4 kDa to 37.1 kDa and changed the maximum number of sialic acids from 14 to 22. As a consequence, carbohydrate content was increased from 40% to 51%. The new carbohydrates were found to be processed in a manner similar to carbohydrates attached to the naturally occurring glycosylation sites and contained no new carbohydrate structures.

When the in vitro bioactivity of darbepoetin alfa was investigated the kinetics and magnitude of EPOR mediated intracellular signaling was similar to rHuEPO, indicating that the additional carbohydrate did not blunt the maximum biological response. However, a slight shift in potency in vitro was observed which correlated with a slightly reduced affinity for EPOR (Figure 2). Studies in normal mice were performed to compare the erythropoietic potencies...
of darbepoetin alfa with that of rHuEPO. In order to elicit a similar erythropoietic response in mice, 3-fold more rHuEPO than darbepoetin alfa was required when dosed at three times per week.74 When weekly dosing paradigms were investigated, 13 fold more rHuEPO was required than darbepoetin alfa for a similar erythropoietic response which increased to approximately 30–40 fold when examined as a single dose.75 Thus the relative in vivo potency of darbepoetin alfa compared to rHuEPO increased as the dosing interval increased.75 Comparative studies were also performed in rodent disease models. In a model of chemotherapy/radiotherapy induced anemia, a 6-fold higher dose of rHuEPO was less effective at stimulating erythropoiesis than darbepoetin alfa.76 Darbepoetin alfa was also found to be effective in treating anemia in a rat model of anemia of inflammation.76 These preclinical studies demonstrated that the further addition of N-linked glycosylation sites could directly increase the in vivo potency of rHuEPO and that darbepoetin alfa was effective in animal models of anemia.

To understand the increase in potency of the rHuEPO analogs, preclinical PK studies were performed. The enhanced glycosylation of darbepoetin alfa was found to increase the serum half-life by 3-fold in rats and dogs when compared to rHuEPO.74 In clinical studies, darbepoetin alfa had an approximately 3-fold longer mean terminal half-life than rHuEPO in patients receiving peritoneal dialysis, with more than a 2-fold greater area under the curve and a 2.5-fold reduced clearance.77 In these patients, the mean half-life of darbepoetin alfa was also found to increase approximately 2-fold when administered subcutaneously compared to intravenous administration.77 Darbepoetin alfa was also found to have an increased serum half-life in anemic patients with nonmyeloid malignancies receiving multiple cycles of chemotherapy when compared with rHuEPO.78 In patients with chemotherapy-induced anemia, this half-life was estimated to be 61–88 hours, depending on timing of chemotherapy.79

One theoretical drawback to the glycoengineering of therapeutic proteins could be immune recognition of the mutated amino acid sequence resulting in the production of neutralizing antibodies. Darbepoetin alfa differs from rHuEPO at five amino acid positions suggesting that an antibody could theoretically be directed against the “altered” region. However, antibody formation was monitored during clinical trials with darbepoetin alfa and no enhanced formation of neutralizing antibodies was observed when compared to epoetin alfa made in the US.79 Indeed, carbohydrates and sialic acid on darbepoetin alfa inhibited antibody binding in immunoassays compared with deglycosylated and desialated molecules.79

**AMG 114 and AMG 205**

Hyperglycosylated analogs of rHuEPO have also been evaluated to determine if further increasing the number of N-linked carbohydrate sites above darbepoetin alfa would further increase the half-life and in vivo potency of the ESA. Analogue AMG 205 was initially generated and contained four additional N-linked glycosylation sites compared to rHuEPO, for a total of seven.80 The increase in glycosylation from two additional sites in darbepoetin alfa to four additional sites in AMG 205, decreased receptor affinity and in vitro biological activity (Table 1 and Figure 2) but increased the serum half-life in rats from 17.2 hours to 31 hours when administered intravenously and 13.2 hours to 28.2 when administered subcutaneously.81 Though the molecular stability of AMG 205 was suitable based on antibody binding assays to an internal epitope recognized by the 9G8a antibody, the O-linked site was not efficiently glycosylated and development of this molecule was discontinued.82

Another hyperglycosylated analog AMG 114, also contained four additional N-linked glycosylation sites but was efficiently glycosylated at the O-linked site and also maintained excellent stability as determined by the 9G8a antibody binding assay.80 This molecule therefore
contained a maximum of 30 sialic acids. The inclusion of four additional N-linked carbohydrates increased the molecule weight from 30.4 kDa in rHuEPO to 44.8 kDa in AMG 114. Molecular structural analyses identified that the additional carbohydrates were distal to the receptor binding sites thus retaining receptor binding activity though AMG 114 also had reduced in vitro activity and receptor affinity compared with rHuEPO and darbepoetin alfa (Table 1 and Figure 2). In dog PK studies, AMG 114 had a serum half-life of 68.9 hours compared to 20.4 hours of darbepoetin alfa. Furthermore, in clinical trials in chemotherapy induced anemia patients, AMG 114 was found to have a serum half-life of approximately 130 hours compared to 73.7 hours with darbepoetin alfa. Dose dependent increases in hemoglobin were observed though modest, and the molecule appeared to be well tolerated. However, AMG 114 was not further developed as darbepoetin alfa was found to meet the existing clinical needs with a synchronous chemotherapy administration schedule.

**Pegylation of rHuEPO**

Several approaches have been taken to generate ESAs with enhanced serum half-life and in vivo bioactivity through the synthetic chemical addition of polyethylene glycol to fully glycosylated rHuEPO.

**Pegzyrepoetin alfa**

Recently, a molecule consisting of a pegylated and fully glycosylated rHuEPO was approved for the treatment of anemia associated with chronic kidney disease (CKD) in several countries. Pegzyrepoetin alfa has been reported to be generated by the addition of methoxy-polyethylene glycol (PEG) polymer chain to the rHuEPO N-terminal amino acid or the ε-amino group on lysines 45 or 52 with a single succinimidyl butanoic acid linker. The PEG used in the conjugation to fully glycosylated rHuEPO is a 30 kDa chain polymer and increases the molecular weight of the molecule to approximately 60 kDa. In preclinical studies, though HhuEPO was more potent than pegzyrepoetin alfa in the induction of cellular proliferation in an EPO dependent cell line UT-7/EPO in vitro, pegzyrepoetin alfa was able to increase reticulocyte count more effectively than identical peptide amounts of rHuEPO in normal mice. In these studies, a single injection of 20 µg/kg of pegzyrepoetin alfa in mice was reported to increase reticulocyte count by 13% vs 7.8% with a comparable peptide dose of rHuEPO. In humans, the PK has been investigated and the serum half-life via both intravenously and subcutaneous administration was approximately 130 hours. In clinical efficacy studies, pegzyrepoetin alfa has been suggested to produce stable hemoglobin levels in CKD patients with once monthly administration. It is currently unclear how pegzyrepoetin alfa is cleared and the mechanisms that account for the enhanced in vivo activity. As with other PEGylated molecules (eg, pegfilgrastim), the attachment of a highly hydrophilic PEG molecule increases the molecular size and therefore reduces renal clearance and extends serum half-life. However, the pegylation may interfere with receptor binding due to steric hindrance and indeed pegzyrepoetin alfa has been reported to have a lower receptor affinity for EPOR.

Recently, a study in sheep to evaluate PK and receptor interactions of HhuEPO and pegzyrepoetin alfa reported pegzyrepoetin alfa was less able to induce EPOR expression in the erythroid compartment in vivo, had slower binding to EPOR and had a reduced internalization and subsequent degradation of surface bound pegzyrepoetin alfa. However, the mechanism for improved half-life and reduced clearance need further elucidation.

**MK-2578**

Another pegylated version of glycosylated HhuEPO, MK-2578, was reported to have reached the clinical development stage. In contrast to Chinese Hamster Ovary synthesis of HhuEPO, the glycosylated HhuEPO in MK-2578 was reported to be synthesized in yeast. Yeast *Pichia pastoris* were genetically engineered to secrete human glycoproteins with fully complex terminally sialylated N-glycans. The yeast cell lines were reported to produce complex glycoproteins with greater than 90% terminal sialylation and were used to synthesize functional HhuEPO that dose dependently increased hematocrit in mice. Though MK-2578 was in clinical development for the treatment of anemia associated with chronic renal insufficiency, this drug was reportedly halted in Phase II trials in 2010 due to an US Food and Drug Administration (FDA) request for cardiovascular outcomes in clinical trials.

**EPO mimetic proteins**

The potential for EPO mimetic proteins to bind, dimerize, and activate the EPOR have been investigated as new modalities to generate ESAs. Several approaches have been taken to evaluate the erythropoietic activity of these agents and improve their half-life and in vivo potency through identification of EPO mimetic peptides (EMP), dimerization of EMPs though covalent chemical approaches and pegylation,
conjugation of EMPs to antibody Fc regions, and generation of agonistic anti-EPOR antibodies.

**EMP1**

Approaches to identify EMP sequences initially used a filamentous phage display library to screen binding phage to the extracellular domain of human EPOR. A 14-mer cyclic peptide with internal disulphide bonds, AF11154, was identified that was able to bind and displace radioactive rHuEPO binding to EPOR, though at a low potency of 10 µM IC₅₀.⁹⁻³-subsequent screening of mutated peptides derived from AF11154 identified peptides including EMP1 that were able to displace radioactive rHuEPO binding with a 50% inhibitory concentration (IC₅₀) of 0.3 µM.⁹⁻³ EMP1 was found to have in vitro bioactivity using EPOR expressing cell line proliferation assays and in vitro erythroid colony formation potential from human bone marrow though the potency was ~9,000-fold and ~40,000-fold less potent than rHuEPO in these assays, respectively.⁹⁻³ The low potency of the EMP1 molecule in vitro also translated to erythropoietic assays in vivo with approximately a 650,000 fold reduced in vivo activity of EMP1 compared to rHuEPO when compared on a molar basis.⁹⁻³,⁹⁻⁴ Pegylated versions of EMP1 dimers were also generated through the addition of the PEG molecules to the N-terminal amino acids of the peptides. This modification increased the potency of the molecule ~50-fold in vitro and ~14-fold in vivo.⁹⁻⁴ Chemical covalent dimerization of the EMP1 peptide was also performed and this was found to increase the potency of the chemical-EMP1 dimer ~100-fold versus the native peptide using in vitro and in vivo assays.⁹⁻⁴,⁹⁻⁵ However, this chemical EMP1 dimer was still reported to be ~6,000-fold less potent than rHuEPO on a molar basis using in vivo erythropoietic assays.⁹⁻⁴,⁹⁻⁵

**CNTO 528 and CNTO 530**

Two approaches using the EMP1 peptide as the EPOR binding domain but fused to Fc proteins have been described to attempt to create an EPOR agonist therapeutic with an extended serum half-life and in vivo biological activity. CNTO 528 has been described to be a fusion protein of EMP1 with the Fc region of human IgG1. In a phase 1 clinical trial in adult healthy volunteers, CNTO 528 was reported to be well tolerated and effective at raising and maintaining hemoglobin levels by at least 1 g/dL following a single intravenous administration.⁹⁻⁶ After a single intravenous administration of CNTO 528, the serum half-life was reported to range from 1.55 to 7.6 days at doses between 0.03 and 0.9 mg/kg.⁹⁻⁶ CNTO 530 is a 58 kDa glycoprotein with two EMP-1 sequences as a pharmacophore and is fused to the human IgG4 Fc domain.⁹⁻⁷ Preclinical studies reported the CNTO 530 competitively bound to the EPOR in UT-7/EPO cells with an IC₅₀ of 17 pM and induced EPOR dependent signaling in the same cell line.⁹⁻⁷ In pharmacokinetic studies in mice the half life (T₁/₂) was ~40 hours and a single dose of 0.3 mg/kg significantly increased erythropoietic parameters (hemoglobin and RBC) that were maintained for ~28 days. More recent studies have reported that CNTO 530 can treat chemotherapy induced anemia in mouse models with 5-fluorouracil or carboplatin plus pachitaxel.⁹⁻⁸ These studies have also suggested that CNTO 530 can provide a sustained activation of ribosomal S6 kinases p70S6K (presumably downstream of EPOR) that was not seen with rHuEPO or darbepoetin alfa, and can also provide an enhanced expansion of bone marrow resident erythroid progenitor pool, though it is unclear how this may occur mechanistically.⁹⁻⁸ The clinical utility of this molecule is however yet to be investigated.

**Peginesatide**

Peginesatide is also a pegylated EMP and recently completed Phase III clinical trials that evaluated the potential of the molecule to treat anemia in patients with chronic renal failure. The preclinical properties of peginesatide have been evaluated and have been reported to compete with native rHuEPO for EPOR binding and displace radioactive EPO from the EPOR with an IC₅₀ ~37 pM versus 7 pM for rHuEPO.⁹⁻⁸ Peginesatide was also reported to induce downstream EPOR signaling through JAK2, STAT5, and AKT in an EPO dependent cell line UT-7/EPO, and induced cellular proliferation in vitro at an EC₅₀ of 460 pM versus 36 pM rHuEPO.⁹⁻⁹,⁹⁻¹⁰ The potential of peginesatide to induce human erythroid colony formation was also evaluated in vitro and was found to be ~16-fold less potent than rHuEPO on a molar basis.⁹⁻¹⁰ In preclinical pharmacokinetic studies, after a single bolus injection of peginesatide, the T½ was reported to be ~22–31 hours in rats and 14–59 hours in monkeys, depending on the dose used with renal elimination as one of the routes of clearance. The potential for peginesatide to enhance erythropoiesis was evaluated in vivo by examining the elevation of reticulocytes and hemoglobin after single intravenous doses between 0.1 and 10 mg/kg in rats and monkeys. In these studies, peginesatide dose dependently increased reticulocytes and induced cellular proliferation in vitro at an EC₅₀ of 100–10² pM. These studies reported the CNTO 530 competitively bound to the EPOR in UT-7/EPO cells with an IC₅₀ of 17 pM and induced EPOR dependent signaling in the same cell line.⁹⁻⁷ In pharmacokinetic studies in mice the half life (T₁/₂) was ~40 hours and a single dose of 0.3 mg/kg significantly increased erythropoietic parameters (hemoglobin and RBC) that were maintained for ~28 days. More recent studies have reported that CNTO 530 can treat chemotherapy induced anemia in mouse models with 5-fluorouracil or carboplatin plus pachitaxel.⁹⁻⁸ These studies have also suggested that CNTO 530 can provide a sustained activation of ribosomal S6 kinases p70S6K (presumably downstream of EPOR) that was not seen with rHuEPO or darbepoetin alfa, and can also provide an enhanced expansion of bone marrow resident erythroid progenitor pool, though it is unclear how this may occur mechanistically.⁹⁻⁸ The clinical utility of this molecule is however yet to be investigated.
application was filed in the USA in May 2011 for the treatment of anemic chronic renal insufficient patients on dialysis and was approved in March 2012. However, an increase in cardiovascular events was observed in CKD patients not on dialysis treated with peginesatide and was not approved for that indication. In addition, due to recent reports of severe allergic reactions including some deaths, the drug was voluntarily recalled in February 2013 and an investigation is ongoing to understand the nature of the adverse reactions to the drug.

One potential theoretical advantage of having a molecule that is immunologically distinct from EPO is that neutralizing antibodies potentially generated to peginesatide in patients would not interact with endogenous EPO, therefore potentially preventing PRCA. This concept was explored in a rat model of PRCA and it was reported that rat anti-EPO antibodies did not cross react with peginesatide and the anemia induced could be treated in the rats. In a clinical study, CKD patients with PRCA were treated with peginesatide and in some patients reticulocytes increased, along with an increase in hemoglobin and decrease in transfusion dependence. However, in this study one patient that had initially responded subsequently developed neutralizing antibodies to peginesatide and became transfusion dependent again.

**Anti-EPOR agonistic antibodies**

Due to the general long in vivo half-life of antibodies, several different approaches have been taken to generate agonistic anti-EPOR antibodies that may provide extended erythropoiesis with less frequent dosing than rHuEPO. The first reported description of agonistic anti-EPO antibodies was by Elliott et al 1996 who generated mouse monoclonal antibodies to the human extracellular region of EPOR. Several of the bivalent antibodies were able to compete with the binding of radioactive EPO to erythroleukemic cell lines, were able to induce the proliferation of EPO dependent UT-7/EPO cells and were able to promote the differentiation of human CD34+ hematopoietic progenitors to erythroid colonies. However, the most potent monoclonal antibody (mAb) mAb 71 was 2–3 log fold less potent than rHuEPO in proliferation assays and 5–6 log fold less potent in the erythroid colony forming assays. However, monovalent Fab fragments of mAb 71 failed to activate the receptor as EPOR requires dimerization and subsequent conformational change to occur for functional activation of signaling. Similar low potency in vitro activity was reported with another anti-EPOR mouse monoclonal antibody. Recently a fully human agonistic anti-EPOR antibody ABT007 was reported to have been generated using the XenoMouse technology. Using in vitro studies, ABT007 was 10-fold less potent using in vitro cell line proliferation assays and 5–40 more ABT007 was needed than rHuEPO to reach a maximum erythroid colony formation from human bone marrow CD34+ hematopoietic progenitors. However, using in vivo studies ABT007 increased hematocrit with once monthly dosing and was reported to be comparable to darbepoetin alfa dosed every 2 weeks in mice. As the antibody was reported not to bind to mouse EPOR, the investigators had to use mice with the humanized EPOR to perform their studies. Interestingly, analysis of the antibodies from these studies found that in vitro and in vivo potency correlated inversely with binding affinity. The development status of ABT007 is at this point unknown.

Recently an approach to generate heterodimeric bi-specific antibodies using combinatorial antibody libraries has been investigated which identified bi-specific antibodies that asymmetrically bind to EPOR to activate JAK/STAT5 pathways and generate hemoglobinized erythroid colonies from human CD34+ progenitors. However, these prototype molecules are still −5-fold less potent than rHuEPO in vitro assays and the in vivo potency at this time is unknown.

**Other approaches**

Several other approaches have been taken to develop ESAs with improved serum half-lives and in vivo erythropoietic activity. In one study, investigators cross-linked cysteine residues in rHuEPO monomers to form rHuEPO dimers and trimers using a chemical modification process. In these studies, the investigators reported that the chemically induced rHuEPO dimers had an extended plasma half-life in comparison with rHuEPO monomers (>24 hours versus 4 hours) in intravenous-injected rabbits, and correlated with an enhanced ability to increase hematocrit in mice (~56% versus ~51%). In a similar study by the same group, the investigators synthesized recombinant rHuEPO dimers joined by a 17 amino acid flexible peptide in COS cells, an African green monkey cell line. Using an in vitro bioassay, the rHuEPO dimers were reported to have an ~5–6-fold increase in specific activity when compared on a molar basis and were reported to have an improved serum half-life and elevated hematocrit above that of rHuEPO alone after a single intravenous dose.

rHuEPO has also been fused to the human neonatal antibody FcRn region of IgG1 to extend the serum half-life and also provide a transport mechanism across pulmonary epithelium as an alternative delivery route. Several molecules were tested including a construct containing two EPO molecules fused to two chains of the FcRn and a chimeric
molecule containing only a single EPO attached to the FcRn dimer. In studies using non-human primates and using an aerosol method of delivery to the lung epithelium, the EPO-Fc monomer more improved PK properties and in vivo erythropoietic responses than the EPO-Fc dimer.

In a phase I clinical trial in normal healthy volunteers, the investigators reported that aerosolized EPO-Fc was able to be delivered by inhalation with dose dependent increases in PK and pharmacodynamic effects. However the current development status of this molecule is unknown.

Conclusions

ESAs have been used successfully to treat anemia in millions of patients, reducing the number of transfusions required. Since the discovery that rHuEPO could effectively increase hemoglobin levels in patients and reduce the numbers of transfusions, alternative approaches have been taken to improve upon the PK and pharmacodynamic properties of new therapeutic molecules. These approaches have included the glycoengineering and PEGylation of rHuEPO and the identification of EPO mimetic proteins conjugated to PEG or fused to antibody Fc regions (Figure 3). In general, these new ESAs share many similarities: the molecules activate the EPOR in the same way and the enhanced erythropoietic activity of the molecules in vivo correlate with extended serum half-life, but the enhanced in vivo activity is inversely related to the in vitro activity and receptor affinity. It will be interesting to determine if newly developed ESAs have the same general properties of existing ESAs or whether alternative approaches to synergize with other mechanisms/receptors to enhance erythropoiesis are efficacious and are developed clinically.

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AMS (author), WS, SL, and BS (see acknowledgments) are employees of Amgen, Inc and hold stock. SE is a consultant for Amgen, Inc and holds stock. JB holds Amgen, Inc stock. The author has no other disclosures to declare.

References


