Epoetin zeta in the management of anemia associated with chronic kidney disease, differential pharmacology and clinical utility

Abstract: Epoetin zeta was granted marketing authorization in October 2007 by the European Medicines Agency as a recombinant human erythropoietin erythropoiesis-stimulating agent to treat symptomatic anemia of renal origin in adult and pediatric patients on hemodialysis and adults on peritoneal dialysis, as well as for symptomatic renal anemia in adult patients with renal insufficiency not yet on dialysis. Currently, epoetin zeta can be administered either subcutaneously or intravenously to correct for hemoglobin concentrations ≤10 g/dL (62 mmol/L) or with dose adjustment to maintain hemoglobin levels at desired levels not in excess of 12 g/dL (75 mmol/L). This review article focuses on epoetin zeta indications in chronic kidney disease, its use in managing anemia of renal origin, and discusses its pharmacology and clinical utility.

Keywords: biosimilar, chronic kidney disease, epoetin alfa, erythropoiesis, renal anemia, Retacrit®

Introduction
Renal anemia occurs as a common complication of chronic kidney disease (CKD). CKD is a complex disease characterized by impaired renal function. The Kidney Disease Improving Global Outcomes (KDIGO) initiative defines CKD as the presence of structural or functional abnormalities of the kidneys resulting in kidney damage, for instance, pathologic abnormalities or markers of kidney damage to include proteinuria, renal tubular syndromes or imaging abnormalities, or level of kidney function measured by glomerular filtration rate (GFR) <60 mL/minute/1.73 m² lasting ≥3 months. There are five stages to disease progression based on estimated GFR levels calculated from serum creatinine levels and levels of proteinuria. These stages range from kidney damage with normal or increased GFR (stage 1) to kidney failure (stage 5). Table 1 shows the KDIGO classifications for the five stages of CKD. The term CKD refers to the presence of any stage of CKD (stage 1 through 5) with or without kidney transplant, and includes both nondialysis and dialysis dependent disease.

Prevalence, disease burden, and treatment costs for CKD are increasing in the US and globally. Overall, in the US, an estimated one in ten adults, or about 20 million individuals, have CKD, and, during the years 1999–2004, data show a higher prevalence among females than males (females 15.0% versus males 11.1%). Although, in 2011, new cases of end stage renal disease (ESRD) declined for the first time in 30 years in the US, there was an overall increase in patients receiving treatment for ESRD. In the UK, CKD affects an estimated 6% of the population. Little data exist for European prevalence. The Global Burden of Disease project’s ranking of leading causes of disability-adjusted...
life years (DALYs) for 291 specific diseases ranks CKD 29th overall globally with regional geographical rankings ranging from 8–44 (see Figure 1). Between 1990 and 2010, DALYs (per 100,000) for CKD show an overall increase of 16.7%, with an even greater percent increase for CKD due to diabetes mellitus (36.1%) or hypertension (42.2%).

Risk factors for developing CKD include age, gender, race, diabetes, and genetic makeup, as well as modifiable factors such as hypertension, proteinuria, anemia, metabolic disturbances, and dyslipidemia. Disease progression to CKD increases the risk for cardiovascular disease, hospitalization, and death, with some suggesting an increased risk for a cardiorenal syndrome.

Hemoglobin (Hb) levels often gradually decline with the decline in renal function. The prevalence of anemia increases as kidney function declines. Renal anemia is associated with adverse patient outcomes, including decreased exercise capacity and quality of life, and increased hospitalization, cardiovascular events, and chance of death. Renal anemia arises from CKD-induced oxidative stress, inflammation, and a relative deficiency in the renal production of erythropoietin (EPO) due to loss of EPO synthesis or inhibitors of EPO. EPO is an endogenous protein produced in the kidneys to stimulate red blood cell production under hypoxic conditions. Similar to the anemia of
chronic disease, renal anemia is normochromic, normocytic, and characteristically hypoproliferative, however different from anemia of chronic disease, renal anemia also shows low EPO and some iron deficiency.\textsuperscript{1,15}

The definition of renal anemia has evolved over the years. Currently, the KDIGO, the European Renal Best Practice group (ERBP), and the National Kidney Foundation (NKF) Kidney Disease Outcomes Quality Initiative (KDOQI) give Hb levels to serve as a guide when diagnosing anemia. The current KDIGO guidelines use the World Health Organization definition of anemia while the ERBP position statement takes into consideration Hb set point differences for European populations.\textsuperscript{30} In the US, the NKF KDOQI continues to use the 2006 KDOQI definition.\textsuperscript{31} Table 2 shows the differences between guidelines for diagnosing renal anemia among NKF KDOQI 2006 US guidelines, KDIGO updated 2012 guidelines, and the ERBP 2013 position statement.\textsuperscript{30,31}

After ruling out other causes of symptomatic anemia in the presence of CKD, and once renal anemia has been diagnosed, renal anemia treatment guidelines suggest a trial of iron therapy before beginning erythropoiesis-stimulating agent (ESA) therapy and based on transferrin saturation (TSAT) and ferritin levels. Iron supplementation improves Hb levels and cellular response to ESA stimulation, which may in turn help reduce the ESA dosage. This ESA response to iron supplementation may arise from how the hepcidin-ferroportin axis helps regulate iron homeostasis by controlling the entry of iron from dietary sources into the bloodstream. The iron mediated hepcidin pathway is suppressed in the presence of hypoxia and anemia, while ESA administration causes hepcidin excess. Hepcidin excess impedes the absorption of dietary iron and the release of iron stores. Therefore, administering iron supplementation, either intravenously for hemodialysis patients, or through an oral agent for nondialyzed CKD patients, for TSAT <30% and ferritin level <300 ng/mL is recommended. Iron supplementation should be used with caution for TSAT >30% and ferritin levels exceeding 500 ng/mL.\textsuperscript{30,32}

In 1988 in Europe, and subsequently for the year 1989 in the US, recombinant human erythropoietin (rHuEPO) was introduced to treat renal anemia. Its ability to normalize Hb led to rHuEPO eventually replacing blood transfusions and the use of androgenic steroids as the mainstay treatment for renal anemia. Moreover, renal anemia was independently associated with the development of left ventricular hypertrophy (LVH) and heart failure. Normalizing Hb with epoetin alfa arrested the progression of LVH in CKD with associated improvements in cardiovascular-related morbidity and mortality.\textsuperscript{37} Additionally, early rHuEPO research reported improved quality of life scores in CKD contributing to the eventual widespread use of rHuEPO in treating renal anemia.\textsuperscript{34–37} Over time, various epoetin analogues (epoetin alfa, epoetin beta, darbepoetin alfa) were developed that effectively increased Hb levels. Patent expirations for rHuEPO biopharmaceuticals led to the development of biosimilars, including epoetin zeta.\textsuperscript{38}

Currently, epoetin zeta carries indications for treating symptomatic renal anemia, for intravenous (IV) and subcutaneous administration in adults and pediatric patients undergoing hemodialysis and in adult patients on peritoneal dialysis, or for those not yet undergoing dialysis.\textsuperscript{39,40} Contraindications include hypersensitivity, patients who develop epoetin-induced antibody-mediated pure red cell aplasia (PRCA) following treatment, uncontrolled hypertension, and patients who for any reason cannot receive adequate antithrombotic prophylaxis.\textsuperscript{39} The purpose of this article is to review the differential pharmacology and clinical utility of epoetin zeta.

### Table 2 Guidelines for diagnosing renal anemia and making further evaluation based on Hb levels

<table>
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<tbody>
<tr>
<td>Adult males and children &gt;15 years</td>
<td>Hb &lt;13.0 g/dL</td>
<td>Hb &lt;13.5 g/dL</td>
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<tr>
<td>Adult females</td>
<td>Hb &lt;12.0 g/dL</td>
<td>Adult males &gt;70 years</td>
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<tr>
<td>Adult females, all ages</td>
<td>Hb &lt;12.0 g/dL</td>
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**Abbreviations:** ERBP, European Renal Best Practice group; Hb, hemoglobin; KDIGO, Kidney Disease Improving Global Outcomes; NKF KDOQI, National Kidney Foundation Kidney Disease Outcomes Quality Initiative; TSAT, transferrin saturation.
OR EPO zeta OR erythropoietin zeta OR Retacrit OR Silapro) AND (kidney* OR renal OR uremia OR uremic OR ESRD OR ESRF OR CRD OR CRF OR CKD) AND (chronic OR insufficient* OR sufficient* OR disease OR fail* OR damage* OR irreversible OR end stage* OR anemia OR anemic). Searches were conducted in PubMed (publication n=10), Embase (publication n=61), Cochrane CENTRAL (publication n=4), Web of Science (publication n=24), ClinicalTrials.gov (publication n=3), and Google Scholar (publication n=278). Additionally, we gathered information from regulatory authorities including the European Medicines Agency (EMA), US Food and Drug Administration (FDA), and US Centers for Medicare and Medicaid Services (CMS), as well as from the professional association The National Kidney Foundation, Inc. Additional pertinent references were identified from the bibliographies of retrieved publications. Finally, we excluded non-English language publications and duplicates.

**Background**

Patent expirations in 2004 for epoetin alfa and 2005 for epoetin beta in the European Union led to the development of biosimilar epoetin. The term “biosimilar”, as used in Europe (termed “follow-on-biologics” in the US and Japan, and called “subsequent-entry biologic” in Canada), refers to officially-approved subsequent versions of innovator biotechnological products with the same primary amino acid sequence as the referent product used at the same dose to treat the same disease.\(^24,28,41,42\) Biosimilars differ from low molecular weight pharmaceutical generics in respect to the size and complexity of the active substance, the heterogeneity of the originator materials, and variability in manufacturing processes.\(^38\)

Pharmaceutical generics are low molecular weight formulations with similar pharmacokinetic and pharmacodynamic profiles as the original that are synthesized from commercially available reagents using standardized procedures.\(^43\)

Biotechnologically derived proteins have highly complex large molecular structures and a high degree of heterogeneity. This creates challenging formulation and manufacturing production processes that prevent the formulation and manufacture of exact copies of the original.\(^44,45\) Unlike generics with commercially available reagents, the biotechnologically derived biosimilars engineer their cellular clones in-house using living organisms or extracted tissues.\(^44-47\) Further, both originator biotechnologicals and biosimilars can undergo changes such as glycosylation or amino acid sequence variation (contamination), among others, that can induce immune effects.\(^43,48-50\) PRCA, an anti-EPO antibody-associated severe anemia that results in transfusion dependence, is one such immune effect.\(^50,51\) Complex manufacturing processes can influence the chemical and clinical characteristics of a biosimilar or affect biosimilar quality. The complex molecular structure, as well as other issues including formulation and manufacture, results in biosimilars being similar to the referent product but not identical. However, biosimilars demonstrate comparable pharmacokinetics and therapeutic equivalence to the reference product.\(^33,45,52-54\) Minor differences in the microheterogeneity pattern of the molecule between the reference product and biosimilar are acceptable only when the difference does not impact safety and efficacy.\(^55\)

Regulatory approval designed to ensure therapeutic equivalence for biosimilars differs from that for generic drug formulations and addresses the complex formulation and manufacture processes. Since the formulation and manufacturing process can affect molecular similarity, regulators face challenges in establishing efficient and appropriate regulatory review and approval pathways to ensure equivalent therapeutic efficacy and safety.\(^57\) The EMA provides regulatory oversight for biosimilars for the EU, with Australia also adopting the EMA comparability and quality guidelines.\(^56\) The EMA guidelines require comparability studies between an authorized originator (termed reference product) biotechnologically derived product and the biosimilar in regards to clinical and nonclinical quality, safety, efficacy,\(^34,45,53,57-60\) and delineate standards for immunogenicity assessment.\(^58,61,62\) Further, the

<table>
<thead>
<tr>
<th>Substance INN</th>
<th>Substance name</th>
<th>Cell line</th>
<th>Brand name</th>
<th>Manufacturer</th>
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<td>Epoetin zeta</td>
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<td>CHO-Zellen</td>
<td>Epobel(^\circledast)</td>
<td>STADA Arzneimittel AG</td>
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<td>Retacrit(^\circledast)</td>
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<td>for distribution in Europe</td>
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**Table 3** Epoetin zeta, substance, cell line, brand name and manufacturer

**Abbreviations:** INN, International nonproprietary names.
EMAn provides specific epoetin guidelines for recombinant EPO biosimilars. In the US, the Biosimilar Price Competition and Innovation Act, passed as part of the 2010 Patient Protection and Affordable Care Act, establishes the framework for the FDA regulatory pathway for biosimilars. To date, the FDA guidance is formative, with the fourth guidance draft released in the spring of 2013. The EMA and FDA have a biosimilar cluster to enhance global development. Also in the US, the CMS issues rules and regulations for physician fee schedule and other Medicare Part B payment systems for biosimilar biological products, as well as billing procedures and other policies affecting the use of EPO stimulating agents in the US Medicare and Medicaid population.

**Pharmacology**

Epoetin zeta is a rHuEPO biosimilar preparation to epoetin alfa. In adults, endogenous EPO production occurs primarily in interstitial kidney cells, as well as in smaller amounts in the liver and central nervous system. Although anemia associated with renal impairment is multifactorial, in the presence of hypoxia, endogenous EPO production declines. Several theories give rationale explaining why hypoxic conditions in renal disease result in reduced EPO production instead of the normal increase in EPO production. Some theories suggest dysfunctional hypoxic cell signaling, whereas others suggest shortened life span of circulating red blood cells, nutritional deficiencies, or inflammation. Dysfunctional hypoxic cell signaling in the kidneys appears to play an important role in the lack of upregulation of EPO in CKD patients. Normally, tissue hypoxia increases the expression of hypoxia inducible transcription factors that bind to a hypoxia-responsive portion of the EPO gene that, in turn, results in increased EPO production. In CKD, the normal oxygen dependent hydroxylation of hypoxia-inducible transcription factors are impaired. Instead of increasing EPO expression, there is an inhibition of EPO production or loss of synthesis of EPO with no corresponding increase in red blood cell production.

Iron deficiency also contributes to renal anemia due to insufficient iron and chronic inflammation. Iron facilitates new red blood cell production whereas iron insufficiency limits the quantity of new red blood cells. Chronic inflammation reduces the amount of iron released from storage when it is needed. Supplemeniting EPO treatments with iron reduces the risk for iron deficiency. Epoetin zeta is a rHuEPO therapy that can be supplemented with iron.

Epoetin zeta is a derivative of the endogenous protein erythropoietin and is a biosimilar of the rHuEPO epoetin alfa, the first commercialized rHuEPO (Epogen®, AMGEN Inc., Thousand Oaks, CA, USA; Procrit®, JANSSEN Biotech, Inc., Horsham, PA, USA; Eprex®, JANSEEN-CILAG Ltd., Buckinghamshire, UK; Epoetin®, AMGEN Inc., Thousand Oaks, CA, USA). As a biosimilar to epoetin alfa, epoetin zeta has the same 165-amino-acid structure and comparable carbohydrate composition with only minor differences in the glycosylation pattern. Glycosylation patterns largely influence immunogenicity and half-life. More complex glycosylation patterns have a higher number of sialic acid residues with longer half-lives. Sialic acid carbohydrate content affects serum half-life, with increased sialic acid content resulting in longer half-life. Epoetin alfa and zeta both have 14 sialic acid residues, and are short acting drugs with a half-life of 6–8 hours when administered intravenously or 19–24 hours if administered subcutaneously. The short half-life allows epoetin zeta to be administered up to three times per week to treat anemia due to CKD.

Epoetin alfa and epoetin zeta are pharmacodynamically comparable both in vitro and in vivo tests. Pharmacokinetic studies also report bioequivalence. Krivoshiev et al report the pharmacokinetics of epoetin zeta and epoetin alfa (the reference product) in healthy volunteers after intravenous and subcutaneous administration, respectively. The pharmacokinetic study assessing intravenous administration was a monocentric, open, randomized, single dose, and two-period crossover trial in 24 volunteers with 21 volunteers included for statistical analysis. Compared to epoetin alfa, the biosimilar epoetin zeta showed nearly identical pharmacokinetic parameters including area under the concentration/time curve, maximal concentration after administration, volume of distribution, total plasma clearance of drug after administration, mean residence time, and terminal half-life. A second study assessing subcutaneous administration of epoetin zeta reported results for a monocentric, double-blind, randomized, single dose, and two-period crossover trial in 48 volunteers. This study evaluated the subcutaneous bioavailability and pharmacokinetic properties of epoetin zeta compared with epoetin alfa (reference product). The subcutaneous bioavailability of epoetin zeta is 24%, which is equivalent to epoetin alfa 20%. Additionally, no significant differences in the aforementioned pharmacokinetic parameters were observed between subcutaneous epoetin zeta and epoetin alfa. The nearly identical pharmacokinetic characteristics suggest bioequivalence of epoetin zeta to epoetin alfa.

The dosage for epoetin zeta is 93–97 IU/kg/week to treat anemia due to loss of endogenous production of EPO in patients with chronic renal failure. Despite its differences in the glycosylation pattern from epoetin alfa, epoetin zeta
stimates erythropoiesis by binding to the EPO receptor on
cells. Once epoetin zeta is bound to the receptor, the receptor
dimerizes and activates the Janus kinase 2 signal pathway,
which goes on to activate the transcription 5 pathway. This
pathway continues to activate the EPO gene needed for
proliferation and maturation of red blood cells and to treat
anemia. As a biosimilar using the endogenous pathway,
there is concern that epoetin zeta could lead to neutral-
izing antibodies. Neutralizing antibodies react against
endoogenous EPO and result in PRCA. PRCA occurs when
there is no endogenous production of EPO due to antibodies
clearing endogenous EPO and rHuEPO. The clearing of all
EPO results in severe anemia requiring blood transfusion
intervention. However, epoetin zeta shows a different immu-
nogenicity profile, mitigating the likelihood of PRCA.

Efficacy
Efficacy studies test the comparability of biosimilars to the
referent product for specific endpoints as well as product
interchangeability. Epoetin zeta demonstrates compar-
able therapeutic efficacy to the reference product (epoetin
alfa) for IV and subcutaneous routes of administration in
the correction and maintenance phases of treating renal
anemia. Hemoglobin endpoints were measured in two premarket
authorization studies for the correction and maintenance
phases of treating renal anemia in patients receiving
hemodialysis. These premarket studies were designed to
assess the clinical efficacy of SB309 (epoetin zeta) to the
referent product, epoetin alfa (Erypo® formulation, Johnson &
Johnson, New Brunswick, NJ, USA). One randomized,
double-blind, verum controlled, multiple dose, parallel group,
multicenter study (study 411-54-04-05-0000) assessed the
therapeutic equivalence of SB309 to the referent product in
the correction phase of renal anemia when administered IV.
Another randomized, double-blind, crossover, verum con-
rolled, multiple dose, multination Phase III trial (study 411-
54-04-04-0000) tested the therapeutic equivalence of SB309
to the referent product in maintaining HB levels, also when
administered IV. Both studies met designated therapeutic
HB endpoints in correcting and maintaining target HB levels
when administered IV compared to the referent product. These
premarket studies, demonstrating a high degree of similarity between
epoetin zeta and the referent product, led to the approval of
epoetin zeta as a biosimilar to epoetin alfa. After receiving
authorization for IV administration, a later premarket
authorization randomized trial assessed the therapeutic
equivalence of subcutaneously administered epoetin zeta
versus epoetin alfa. This study showed therapeutic equiva-
lence in maintaining target HB levels (epoetin zeta mean
HB 10.94 ± 0.84 g/dL, epoetin alfa mean HB 11.02 ± 0.94 g/dL
[95% CI –0.28 g/dL to 0.12 g/dL]) and safety profiles.

Postmarket authorization research reports epoetin zeta
as meeting or maintaining target HB levels when adminis-
tered IV, over the long term, or when patients switch from
other ESAs to epoetin zeta. A study assessing the long-term
safety and tolerability of IV epoetin zeta administration
reports maintaining target HB levels of 10.5–12.5 g/dL (after
56 weeks in n=745 patients, mean HB=11.3–11.5 g/dL; after
108 weeks in n=164 patients, mean HB=11.1–11.6 g/dL)
with stable dosing and no neutralizing antibodies against
erythropoietin. One German postmarketing surveillance
study of IV administered epoetin zeta in 322 eligible elderly
(age ≥75 years) patients from 40 centers showed epoetin
zeta as effective at maintaining HB levels with no significant
difference in mean dose requirements or safety concerns.
A study by Bajraktar et al assessing interchangeability
switched 33 patients in two dialysis centers from various ESAs
to epoetin zeta. The Bajraktar et al study reported that epoetin
zeta maintained target HB with equivalent mean dose, no
PRCA, and a safety profile in line for the study population.

Safety
Several studies report adverse events associated with using
erthropoiesis stimulating agents (ESAs), including epoetin
alfa, when trying to achieve supraphysiologic target HB levels or
when using high doses to achieve normalization of HB levels in
poorly responsive renal anemia. When treating anemia in
CKD with ESAs, using high or supraphysiologic HB thresholds,
and/or high target HB levels, and/or high ESA dosing levels
to guide therapy a significant association is shown, with an
increased risk for mortality, morbidity, cardiovascular events
and stoke, and tumor progression. Also, supraphysiologic HB
thresholds significantly increase the risk for an immunogenic
response termed ESA-induced PRACA.

Three important studies – the Correction of Hemoglobin
and Outcomes in Renal Insufficiency (CHOIR), Cardiovascular
Reduction Early Anemia Treatment Epoetin beta (CREATE),
and Trial to Reduce cardiovascular Events with Aranesp Therapy
(TREAT) – reported pivotal findings influencing current thinking about safety and prac-
tice guidelines related to target HB ranges and dosing for
all ESAs, including biosimilars such as epoetin zeta when
treatment of renal anemia. CHOR and CREATE reported findings from an open-label research design, while the TREAT study reported results from a placebo controlled randomized double-blind controlled trial. The CHOR study (EPO alfa) included 1,432 CKD patients, approximately 50% of whom had coexistent diabetes, and had an intention-to-treat Hb range of 13.5 g/dL versus 11.3 g/dL. The study was stopped due to the number and severity of adverse safety events in the high Hb group.82 Another study, CREATE, assessed EPO beta with Hb intention-to-treat (13–15 g/dL versus 10.5–11.5 g/dL) in 603 patients with CKD and anemia (n=301 and n=302, respectively) with results showing significant progression to ESRD in the higher target Hb group.83 The study also showed no significant difference between groups for cardiovascular outcomes and LVH.83 A third study, TREAT, included patients with CKD, type 2 diabetes, and anemia (n=2,012 treated with darbepoetin alfa targeting Hb 13 g/dL versus the placebo control group n=2,026 treated with rescue doses of darbepoetin alfa for Hb <9 g/dL) and reported no significant difference in hazard rates for cardiovascular events, death, myocardial ischemia, or ESRD, and a significant nearly two-fold increased risk of stroke.84

Current clinical practice recommendations suggest lower than previously accepted parameters for target Hb ranges in CKD for those not yet on dialysis and for patients on dialysis, as well as for low risk and high risk patients with other comorbid conditions. Guidelines continue to evolve in light of the CHOR, CREATE, and TREAT findings and subsequent secondary analysis of their data, among other research using secondary analysis of the data, with revisions being made to previous standards for appropriate Hb levels when initiating treatment as well as target Hb ranges for maintenance therapy. To give context, the 2008 ERBP guidelines for the management of anemia in CKD revised earlier guidelines and emphasize safety concerns when using ESAs.85 More recently, in April 2013, the KDIGO position paper on anemia management guidelines in CKD summarizes, in chapter 3, the recent evidence about earlier EPO safety concerns and recommends adapting the guidelines for the European population.29 When considering the risks and benefits of initiating ESA therapy, the KDIGO recommends caution in high risk groups (stroke, vascular access loss, or hypertension) and great caution for those with CKD and active malignancy, and suggests Hb levels for low-risk patients (no higher than 12 g/DL) and high risk patients (Hb between 9 and 10 g/DL) when initiating therapy.29 The report discusses Hb normalization and suggests individualized therapy, but cautions against using ESAs to maintain Hb levels above 11.5 g/dL.30 The KDIGO group goes on to say ESAs should not be used to intentionally increase Hb concentrations above 13 g/dL (130 g/L).30 In the US, in March 2007, the FDA issued a public health advisory with an updated black box warning for approved ESAs in the US (epoetin alfa and darbepoetin alfa) with the recommendation to monitor Hb levels and adjust ESA dose to the lowest possible to avoid blood transfusion.84 Collaterally, the CMS put forth a position statement for ESA use in the treatment of anemia in adults with CKD, including patients on dialysis and patients not on dialysis, after reviewing the evidence related to safety.55,66

The safety profile for epoetin zeta is similar to epoetin alfa, the reference product. Reports of ESA-induced PRACA in all commercially available ESAs are a key safety concern. Although the mechanism behind ESA-induced PRACA remains unknown, the highest incidence arose from epoetin alfa manufactured outside the US (Eprex®/Erypro®; Ortho-biologics, LLC, Manati, Puerto Rico).86 Our review found no reports for ESA-induced PRACA related to manufacturer site. The Baldamus et al study assessing the long-term safety of IV administered epoetin zeta showed no incidence of PRACA.79 Three other postmarket clinical trials reported no cases of anti-epoetin antibodies or PRCA.54,28,41

A few studies have compared epoetin alfa’s safety profile to that of epoetin zeta. Del Vecchio and Locatelli’s review article examined cardiovascular safety (in particular stroke and hypertension), cancer progression, and immunogenicity associated with using ESA to treat anemia in patients with chronic disease.57 To avoid high ESA dosing, the authors suggest using a conservative and individualized approach based on a risk/benefit evaluation, and targeting intermediate Hb levels.75,86

Eight studies reported the safety of epoetin zeta in patients with renal anemia. Of those, there are four clinical trials,24,28,41,79 three observational studies,76,81,87 and one post hoc analysis based on two clinical trials.50 Table 4 summarizes the common adverse events observed in more than 5% of the patients in the four clinical trials. Infections and infestations were the most common adverse events reported in the clinical trials (12.5%–34.1%) followed by gastrointestinal disorders (5.2%–21.9%) and injury, poisoning, and procedural complications (7.2%–25.8%). Baldamus et al79 reported 715 serious adverse events in 278 patients with cardiac disorders (8.6% of patients), vascular disorders (6.6% of patients), and injury, poisoning, and procedural complications (7.0% of patients) as the most common groups of serious adverse events occurring in more than 5% of patients. Krivosheiev et al41 assessed epoetin zeta administered subcutaneously for
Table 4 Adverse events reported in epoetin zeta clinical trials

<table>
<thead>
<tr>
<th>Study design</th>
<th>Baldamus et al29</th>
<th>Krivosheiev et al41</th>
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<tr>
<td>Study population (n)</td>
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<td>149 (20.0)</td>
<td>17 (7.3)</td>
<td>17 (7.3)</td>
<td>20 (6.6)</td>
</tr>
<tr>
<td>Hypertension</td>
<td>81 (10.9)</td>
<td>27 (11.6)</td>
<td>16 (5.2)</td>
<td>27 (8.6)</td>
</tr>
<tr>
<td>Respiratory, thoracic, and mediastinal disorders</td>
<td>163 (21.9)</td>
<td>27 (11.6)</td>
<td>16 (5.2)</td>
<td>43 (13.7)</td>
</tr>
<tr>
<td>Gastrointestinal disorders</td>
<td>13 (4.2)</td>
<td>13 (4.2)</td>
<td>13 (4.2)</td>
<td>13 (4.2)</td>
</tr>
<tr>
<td>Skin and subcutaneous disorders</td>
<td>105 (14.1)</td>
<td>16 (6.9)</td>
<td>16 (5.2)</td>
<td>27 (8.6)</td>
</tr>
<tr>
<td>Musculoskeletal and connective tissue disorders</td>
<td>78 (10.5)</td>
<td>9 (3.9)</td>
<td>9 (3.9)</td>
<td>22 (7.0)</td>
</tr>
<tr>
<td>General disorders and administration site conditions</td>
<td>192 (25.8)</td>
<td>23 (9.9)</td>
<td>22 (7.2)</td>
<td>35 (11.2)</td>
</tr>
<tr>
<td>Injury, poisoning, and procedural complications</td>
<td>93 (12.5)</td>
<td>15 (6.5)</td>
<td>15 (6.5)</td>
<td>16 (5.1)</td>
</tr>
<tr>
<td>Surgical and medical procedures</td>
<td>278 (37.3)</td>
<td>38 (16.4)</td>
<td>54 (17.7)</td>
<td>43 (13.7)</td>
</tr>
<tr>
<td>Serious adverse events</td>
<td>715</td>
<td>91</td>
<td>71</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: EPO, erythropoietin; PRCA, pure red cell aplasia.

maintenance treatment of renal anemia. The study reported 38 patients as having 91 serious adverse events, with the three most common serious adverse events being medical surgical and medical procedures (4.7% of patients), cardiac disorders (3.4% of patients), and nervous system disorders (3.4%). Krivosheiev et al42 also compared therapeutic effects of epoetin zeta and epoetin alfa in the correction of renal anemia, where 54 out of 305 patients treated with epoetin zeta reported serious events. Wizemann et al24 crossover design clinical trial showed 43 patients experiencing 71 serious adverse events, with injuries, poisoning, and procedural complications as the most reported serious adverse events (23.3% of patients).

Lonnemann and Wrenger’s76 small clinical observational study assessing the use of epoetin zeta in ESRD reported 17 out of 18 patients as switching from various ESAs to epoetin zeta with no side effects after 6 months of epoetin zeta treatment. Bajraktar et al55 evaluated the safety of epoetin zeta in 33 dialysis patients and reported that 3% of the patients develop hypotension, in-dialyzers clotting, or thrombosis of the arteriovenous fistula. Perani et al57 examined the immunological reaction of epoetin zeta in 12 patients with CKD. This study showed no adverse events after switching from methoxy polyethylene glycol epoetin beta to epoetin zeta.

In a post hoc analysis of three large Phase III clinical trials, Wiecek et al58 reported the safety of switching epoetin alfa and epoetin zeta. The post hoc analysis includes 481 patients from the maintenance trial (n=239) as well as from the induction trial (n=242). The results show no difference in the treatment-emergent adverse event profiles when patients switch from epoetin alfa to epoetin zeta. The same type of treatment-emergent events occurred as would be seen in ≥10% of the patient population.

Overall, the biosimilar epoetin zeta shows tolerable adverse effects in patients with renal anemia compared to the reference product epoetin alfa.52,59 However, regulations require a detailed risk management plan to monitor serious and severe safety events and unknown potential safety issues for pharmacovigilance.77 For instance, assessing safety issues in the populations underrepresented in clinical trials, such as elderly patients or patients with various comorbidities, would add to risk management plan requirements. Additional postmarketing pharmacovigilant activities to
evaluate unanticipated issues that may happen when patients switch between a biosimilar and their referent product could include the close monitoring of PRCA and thromboembolic events, as well as drug utilization and tracking of rare but serious adverse events.\textsuperscript{57,89} Table 4 illustrates the adverse events reported in epoetin zeta clinical trials.

**Current treatment in CKD**

Patients with symptomatic renal anemia subsequent to ESRD or CKD of at least stage 3 (estimated GFR <60) whose Hb levels are less than or equal to 10 g/dL are eligible for treatment.

For all CKD indications, current EMA guidelines recommend specific target Hb ranges and dosing regimens for initiating treatment (termed the correction phase) as well as for maintenance therapy. Target Hb levels are particularly important due to the safety concerns surrounding the increased risk for mortality and morbidity associated with epoetin use to achieve relatively high target Hb levels.\textsuperscript{90} The target Hb ranges are 10–12 g/dL (6.2–7.5 mmol/L) for adults and 9.5–11 g/dL (5.9–6.8 mmol/L) in pediatric patients. Clinicians should avoid maintaining Hb levels >12 g/dL (7.5 mmol/L) and should be vigilant to avoid a rise in Hb levels >2 g/dL (1.25 mmol/l) over a 4 week period.\textsuperscript{90}

Licensed health care providers face patient care challenges related to the interchangeability of biotherapeutics and ESA drug safety concerns including those for epoetin zeta.\textsuperscript{91,92} Interchangeability can occur between biosimilar products and the reference product, as well as among a biopharmaceutical product class (eg, any ESAs). Salem and Harvie discuss the nurse’s role and responsibilities regarding biosimilar medicines and their clinical use.\textsuperscript{93} Inadequate knowledge of biosimilar products increases the risks for medication errors and adverse events, and/or may affect desired treatment outcomes. Due to the complex nature of biopharmaceutical products, the study authors suggest providing additional professional training and educational opportunities for healthcare providers who prescribe, dispense, or administer biopharmaceutical products or biosimilars.\textsuperscript{93} Health care providers should be well informed about the differences between biosimilars and their reference products, gain evidence-based knowledge about biosimilar use in clinical practice, and track adverse events when patients switch between biosimilar and reference products.\textsuperscript{78,92,93}

**Pharmacoeconomic considerations**

ESAs bring significant clinical benefits to patients with renal anemia. However, ESAs are very expensive and access to them is likely to be restricted. With the advent of biosimilars, drug access and treatment options improved adding potential health economic benefits over the reference biopharmaceutical ESA therapy.\textsuperscript{87,88,93,94,96} In one survey in Spain, the epoetin zeta cost analysis reported a total cost saving of nearly 45% with respect to epoetin alfa.\textsuperscript{97} Costs per patient per month declined from €298 for epoetin alfa therapy to €177 per patient per month for epoetin zeta.\textsuperscript{96} Additionally, a UK cost minimization study showed that epoetin zeta minimizes the costs for treating renal anemia compared to epoetin alfa.\textsuperscript{94} Epoetin zeta decreased the cost by £7.9 per week for a hemodialysis patient in the Hb correction phase and by £3.9–£15.6 per week in a hemodialysis patient at Hb maintenance phase, based on a 70 kg patient.\textsuperscript{98} Low acquisition costs for epoetin zeta contribute to the potential cost savings. However, the risk of immunogenicity and the safety profile for serious adverse events raises uncertainties for the long-term safety and effectiveness of biosimilars. Future epoetin zeta studies could assess patient outcomes and costs for target Hb levels and further compare results between epoetin zeta and epoetin alfa. Additionally, Simoens suggests conducting further pharmacoeconomic analyses at multiple time points throughout the life cycle of biological pharmaceutical products.\textsuperscript{99}

**Utility**

Utility measures the value of health states from a specified perspective (eg, society, payer, or patient perspective) using a standard measure such as a quality adjusted life year (QALY). Our search did not find any formal cost utility analysis or direct elicitation measures, such as standard gamble or time trade-off, for epoetin zeta in CKD. We found only one study reporting quality of life (QOL) for epoetin zeta in chemotherapy-induced anemia.\textsuperscript{100}

Early research suggests improved health outcomes and symptom management for rHuEPO (EPO alfa), such as exercise capacity,\textsuperscript{36} and health related QOL in patients with renal anemia.\textsuperscript{37,101,102} A recent post hoc analysis of the Canadian Erythropoietin Study Group trial (EPO alfa) included patients on dialysis >3 months with anemia (Hb <9 g/dL), randomized to (n=38, epoetin zeta intravenous treatment group versus 40, placebo), intention-to-treat Hb range 9.5–11 g/dL EPO alfa treatment group versus Hb 11.5–13 g/dL placebo control) showed significantly that fatigue, shortness of breath, and weakness were lessened, and energy was improved in the epoetin alfa treatment group compared to controls.\textsuperscript{103} Another post hoc, multicenter, open-label prospective study using epoetin alfa to treat anemia in patients with nondialysis CKD showed that, across every 2 g/dl change in Hb, the greatest incremental improvement in QOL occurred when the Hb level
reached 11–12 g/dL. A meta-analysis of epoetin alfa clinical trials by Jones et al reported improvements in Hb levels and QOL, and reductions in hospitalizations, overall transfusion rate, and number of units of blood transfused. Another study showed improvements in QOL when treating anemia with epoetin alfa in patients with human immunodeficiency virus infection/acquired immunodeficiency syndrome and coexistent CKD.

However, the CHOIR trial reported serious increased risk of death, myocardial infarction, congestive heart failure, and stroke with no incremental improvement in QOL when correcting renal anemia with epoetin alfa with target Hb levels of 13.5 g/dL. Target Hb levels remain an important consideration in the risk benefit profile for epoetin alfa and in the relationship to utility measures and health outcomes. A review by Leaf and Goldfarb to assess the magnitude and nature of ESA-associated improvements in health related QOL found the maximal increase in health related QOL occurs for Hb ranging between 10 and 12 g/dL, with health-related quality of life (HRQL). improvements blunted beyond that range. Additional utility research is needed to assess QALYs, symptom management using the current standard of care for appropriate Hb levels when initiating therapy, and Hb target ranges to correct renal anemia with epoetin zeta compared to other EPAs or biosimilars.

Although we did not find any studies reporting QOL outcomes generally or in terms of QALY, or by symptom management for epoetin zeta in managing anemia in CKD, one study reported improvements in QOL when treating patients with chemotherapy-induced anemia with epoetin zeta. Future research could further our knowledge about utility outcomes for epoetin zeta in CKD, by the different stages of CKD, or among patients with different risk profiles as compared to the reference product (EPO alfa), or as compared to other ESAs or other biosimilars.

Conclusion

Our review of the biosimilar epoetin zeta found that epoetin zeta has therapeutic equivalence and a comparable safety profile to the reference product (EPO alfa). Epoetin zeta shows comparable efficacy to epoetin alfa in the correction and maintenance phases of CKD anemia treatment for both IV and subcutaneous administration, as well as when interchanged with other ESAs. Although two studies report cost savings, the small number of rigorous studies devoted to cost effectiveness limits the generalizability of suggesting epoetin zeta as a cost effective alternative to current standard treatment or other ESAs. Although one study assessed health related QOL in cancer, to date, we found no utility studies assessing measures of health related QOL, when treating renal anemia with epoetin zeta. Evidence based practice guidelines for ESAs for both originator products and biosimilars, including epoetin zeta, continue to evolve in light of safety concerns, and include recent recommendations about contraindications, when to initiate therapy, dosing, and parameters for Hb levels and target maintenance Hb ranges. Further research is needed to expand our knowledge of the cost effectiveness and health related QOL outcomes for epoetin zeta.

Disclosure

The authors report no conflicts of interest in this work.

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