NEW ZEALAND DATA SHEET

1. PRODUCT NAME
PRAXBIND 50 mg/mL solution for injection/infusion

2. QUALITATIVE AND QUANTITATIVE COMPOSITION
Each mL of solution for injection/infusion contains 50 mg idarucizumab.

Each vial contains 2.5 g idarucizumab in 50 mL.

Idarucizumab is produced by recombinant DNA technology in Chinese Hamster Ovary cells.

Excipients with known effect:
Each 50 mL vial contains 2 g sorbitol and 25 mg sodium (see section 4.4).

For full list of excipients, see section 6.1

3. PHARMACEUTICAL FORM
Solution for injection/infusion.

Clear to slightly opalescent, colourless to slightly yellow solution.

4. CLINICAL PARTICULARS
4.1 Therapeutic indications
Praxbind is a specific reversal agent for dabigatran and is indicated in patients treated with Pradaxa when rapid reversal of the anticoagulant effects of dabigatran is required:
- For emergency surgery/urgent procedures
- In life-threatening or uncontrolled bleeding.

4.2 Dose and method of administration

Dose
The recommended dose of Praxbind is 5 g (2 x 2.5 g/50 mL).

In a limited number of patients, recurrence of plasma concentrations of unbound dabigatran and concomitant prolongation of clotting tests have occurred up to 24 hours after administration of idarucizumab (see section 5.1).

Administration of a second 5 g dose of Praxbind may be considered in the following situations:
- recurrence of clinically relevant bleeding together with prolonged clotting times, or
- patients require a second emergency surgery/urgent procedure and have prolonged clotting times.

Relevant coagulation parameters are activated Partial Thromboplastin Time (aPTT), diluted Thrombin Time (dTT) or Ecarin Clotting Time (ECT) (see section 5.1).

Restarting Antithrombotic Therapy
Pradaxa treatment can be re-initiated 24 hours after administration of Praxbind, if the patient is clinically stable and adequate hemostasis has been achieved.
After administration of Praxbind, other antithrombotic therapy (e.g. low-molecular weight heparin) can be started at any time, if the patient is clinically stable and adequate hemostasis has been achieved.

Absence of antithrombotic therapy exposes patients to the thrombotic risk of their underlying disease or condition.

**Patients with renal impairment**
No dose adjustment is required in renally impaired patients. Renal impairment did not impact the reversal effect of idarucizumab (see section 5.2).

**Patients with hepatic impairment**
No dose adjustment is required in hepatically impaired patients (see section 5.2).

**Elderly**
No dose adjustment is required in elderly patients aged 65 years and above (see section 5.2).

**Paediatric population**
The safety and efficacy of Praxbind in children below the age of 18 years have not yet been established. No data are available.

**Method of administration**
Intravenous use.

Praxbind (2 x 2.5 g/50 mL) is administered intravenously as two consecutive infusions over 5 to 10 minutes each or as a bolus injection.

For additional instructions for use and handling see section 6.6.

**4.3 Contraindications**
None.

**4.4 Special warnings and precautions for use**
Idarucizumab binds specifically to dabigatran and reverses its anticoagulant effect. It will not reverse the effects of other anticoagulants (see section 5.1).

Praxbind treatment can be used in conjunction with standard supportive measures, which should be considered as medically appropriate.

**Traceability**
In order to improve traceability of biological medicinal products, the trade name and the batch number of the administered product should be clearly recorded in the patient file.

**Hypersensitivity**
The risk of using Praxbind in patients with known hypersensitivity (e.g. anaphylactoid reaction) to idarucizumab or to any of the excipients needs to be weighed cautiously against the potential benefit of such an emergency treatment. If an anaphylactic reaction or other serious allergic reaction occurs, administration of Praxbind should be discontinued immediately and appropriate therapy initiated.
Hereditary fructose intolerance

The recommended dose of Praxbind contains 4 g sorbitol as an excipient. In patients with hereditary fructose intolerance, parenteral administration of sorbitol has been associated with reports of hypoglycemia, hypophosphatemia, metabolic acidosis, increase in uric acid, acute liver failure with breakdown of excretory and synthetic function, and death. Therefore, in patients with hereditary fructose intolerance the risk of treatment with Praxbind must be weighed against the potential benefit of such an emergency treatment.

Thromboembolic Events

Patients being treated with dabigatran have underlying disease states that predispose them to thromboembolic events. Reversing dabigatran therapy exposes patients to the thrombotic risk of their underlying disease. To reduce this risk, resumption of anticoagulant therapy should be considered as soon as medically appropriate (see section 4.2).

Sodium

This medicinal product contains 2.2 mmol (or 50 mg) sodium per dose. This should be taken into consideration by patients on a controlled sodium diet.

4.5 Interaction with other medicines and other forms of interaction

No formal interaction studies with Praxbind and other medicinal products have been performed. Based on the pharmacokinetic properties and the high specificity in binding to dabigatran, clinically relevant interactions with other medicinal products are considered unlikely.

Preclinical investigations have shown no interactions with volume expanders, coagulation factor concentrates and anticoagulants other than dabigatran (see section 5.1).

4.6 Fertility, pregnancy and lactation

Pregnancy

There are no data for the use of Praxbind in pregnant women. Reproductive and developmental toxicity studies have not been performed, given the nature and the intended clinical use of the medicinal product. Praxbind may be used during pregnancy, if the expected clinical benefit outweighs the potential risks.

Breast-feeding

It is unknown whether idarucizumab is excreted in human milk.

Fertility

There are no data on the effect of Praxbind on fertility.

4.7 Effects on ability to drive and use machines

Not relevant.

4.8 Undesirable effects

In a phase III trial the safety of Praxbind has been evaluated in 503 patients, who had uncontrolled bleeding or required emergency surgery or procedures and were under treatment with Pradaxa (dabigatran etexilate), as well as in 224 volunteers in phase I trials.
No adverse reactions have been identified.

**Reporting of suspected adverse reactions**
Reporting suspected adverse reactions after authorisation of the medicine is important. It allows continued monitoring of the benefit/risk balance of the medicine. Healthcare professionals are asked to report any suspected adverse reactions https://nzphvc.otago.ac.nz/reporting/

### 4.9 Overdose

There is no clinical experience with overdoses of Praxbind.

The highest dose of Praxbind studied in healthy subjects was 8 g. No safety signals have been identified in this group.

For advice on the management of overdose please contact the National Poisons Centre on 0800 POISON (0800 764766).

### 5. PHARMACOLOGICAL PROPERTIES

#### 5.1 Pharmacodynamic properties

Pharmacotherapeutic group: all other therapeutic products, antidotes  
ATC code: V03AB37

**Mechanism of action**

Idarucizumab is a specific reversal agent for dabigatran. It is a humanised monoclonal antibody fragment (Fab) that binds to dabigatran with very high affinity, approximately 300-fold more potent than the binding affinity of dabigatran for thrombin. The idarucizumab-dabigatran complex is characterised by a rapid on-rate and extremely slow off-rate resulting in a very stable complex. Idarucizumab potently and specifically binds to dabigatran and its metabolites and neutralises their anticoagulant effect.

**Pharmacodynamic effects**

The pharmacodynamics of idarucizumab after administration of dabigatran etexilate were investigated in healthy subjects aged 45 to 64 years receiving a dose of 5 g via intravenous infusion. The median peak dabigatran exposure in the investigated healthy subjects was in the range of a twice daily administration of 150 mg dabigatran etexilate in patients.

*Effect of idarucizumab on the exposure and anticoagulant activity of dabigatran*

Immediately after the administration of idarucizumab, the plasma concentrations of unbound dabigatran were reduced by more than 99%, resulting in levels with no anticoagulant activity.

The majority of the patients showed sustained reversal of dabigatran plasma concentrations up to 12 hours (≥90%). In a subset of patients, recurrence of plasma levels of unbound dabigatran and concomitant elevation of clotting tests was observed, possibly due to redistribution of dabigatran from the periphery. This occurred 1-24 hours after administration of idarucizumab mainly at timepoints ≥12 hours.
Dabigatran prolongs the clotting time of coagulation markers such as diluted Thrombin Time (dTT), Thrombin Time (TT), activated Partial Thromboplastin Time (aPTT) and Ecarin Clotting Time (ECT), which provide an approximate indication of the anticoagulation intensity. A value in the normal range after administration of idarucizumab indicates that a patient is no longer anticoagulated. A value above the normal range may reflect residual active dabigatran or other clinical conditions e.g., presence of other drugs or transfusion coagulopathy. These tests were used to assess the anticoagulant effect of dabigatran. A complete and sustained reversal of dabigatran-induced clotting time prolongation was observed immediately after the idarucizumab infusion, lasting over the entire observation period of at least 24 hours.
Figure 2 – Reversal of dabigatran-induced clotting time prolongation determined by dTT in the representative group of healthy subjects (administration of idarucizumab or placebo at 0 hours)

**Thrombin generation parameters**

Dabigatran exerts pronounced effects on parameters of the endogenous thrombin potential (ETP). Idarucizumab treatment normalised both thrombin lag time ratio and time to peak ratio to baseline levels as determined 0.5 to 12 hours after the end of the idarucizumab infusion. Idarucizumab alone has shown no procoagulant effect measured as ETP. This suggests that idarucizumab has no prothrombotic effect.

**Re-administration of dabigatran etexilate**

24 hours after the idarucizumab infusion, re-administration of dabigatran etexilate resulted in expected anticoagulant activity.

**Immunogenicity**

Serum samples from 283 subjects in phase I trials (224 volunteers treated with idarucizumab) and 501 patients were tested for antibodies to idarucizumab before and after treatment.

Pre-existing antibodies with cross-reactivity to idarucizumab were detected in approximately 12% (33/283) of the phase I subjects and 3.8% (19/501) of the patients. No impact on the pharmacokinetics or the reversal effect of idarucizumab or hypersensitivity reactions were observed.

Treatment-emergent possibly persistent anti-idarucizumab antibodies with low titres were observed in 4% (10/224) of the phase I subjects and 1.6% (8/501) of the patients suggesting a low immunogenic potential of idarucizumab. In a subgroup of 6 phase I subjects, idarucizumab was administered a second time, two months after the first administration. No anti-idarucizumab antibodies were detected in these subjects prior to the second administration. In one subject, treatment-emergent anti-idarucizumab antibodies were detected after the second administration. Nine patients were re-dosed with idarucizumab. All nine patients were re-dosed within 6 days after the first idarucizumab dose. None of the patients re-dosed with idarucizumab tested positive for anti-idarucizumab antibodies.
**Preclinical pharmacodynamics**
A trauma model in pigs was performed using a blunt liver injury after dosing with dabigatran to achieve supratherapeutic concentrations of about 10-fold of human plasma levels. Idarucizumab effectively and rapidly reversed the life-threatening bleeding within 15 minutes after the injection. All pigs survived at idarucizumab doses of approximately 2.5 and 5 g. Without idarucizumab, the mortality in the anticoagulated group was 100%.

Preclinical investigations with idarucizumab have shown no interactions with:
- volume expanders
- coagulation factor concentrates, such as prothrombin complex concentrates (PCCs, e.g. 3 factor and 4 factor), activated PCCs (aPCCs) and recombinant factor VIIa
- other anticoagulants (e.g. thrombin inhibitors other than dabigatran, Factor Xa inhibitors including low-molecular weight heparin, vitamin K-antagonists, heparin). Thus idarucizumab will not reverse the effects of other anticoagulants.

**Clinical efficacy and safety**
Three randomised, double-blind, placebo-controlled Phase I studies in 283 subjects (224 treated with idarucizumab) were conducted to assess the safety, efficacy, tolerability, pharmacokinetics and pharmacodynamics of idarucizumab, given alone or after administration of dabigatran etexilate. The investigated population consisted of healthy subjects and subjects exhibiting specific population characteristics covering age, body weight, race, sex and renal impairment. In these studies the doses of idarucizumab ranged from 20 mg to 8 g and the infusion times ranged from 5 minutes to 1 hour.

Representative values for pharmacokinetic and pharmacodynamics parameters were established on the basis of healthy subjects aged 45-64 years receiving 5 g idarucizumab (see sections 5.1 and 5.2).

One prospective, open-label, non-randomised, uncontrolled study (RE-VERSE AD) was conducted to investigate the treatment of adult patients who presented with dabigatran-related life-threatening or uncontrolled bleeding (Group A) or who required emergency surgery or urgent procedures (Group B). The primary endpoint was the maximum percentage reversal of the anticoagulant effect of dabigatran within 4 hours after the administration of idarucizumab, based on central laboratory determination of dilute thrombin time (dTT) or ecarin clotting time (ECT). A key secondary endpoint was the restoration of haemostasis.

RE-VERSE AD included data for 503 patients: 301 patients with serious bleeding (Group A) and 202 patients requiring an urgent procedure/surgery (Group B). Approximately half of the patients in each group were male. The median age was 78 years and the median creatinine clearance was 52.6 mL/min. 61.5% of patients in Group A and 62.4% of patients in Group B had been treated with dabigatran 110 mg twice daily.

Reversal was only evaluable for those patients showing prolonged coagulation times prior to idarucizumab treatment. Most patients, in both Groups A and B, achieved complete reversal of the anticoagulant effect of dabigatran (dTT: 98.7%; ECT: 82.2%; aPTT: 92.5% of evaluable patients, respectively) in the first 4 hours after administration of 5 g idarucizumab. Reversal effects were evident immediately after administration.
Figure 3 – Reversal of dabigatran-induced clotting time prolongation determined by dTT in patients from the RE-VERSE AD study (N=487)
Figure 4 – Reversal of dabigatran-induced clotting time prolongation determined by ECT in patients from RE-VERSE AD study (N=487)

<table>
<thead>
<tr>
<th>Time post Idarucizumab</th>
<th>ECT [s]</th>
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<tr>
<td>Baseline</td>
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<td>Between 10-30 min vials</td>
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<td>12 h</td>
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<td>24 h</td>
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Individual data with median and 25th/75th percentiles
10th/90th percentiles
5th/95th percentiles
Upper limit normal
Figure 5 – Reversal of dabigatran-induced clotting time prolongation determined by aPTT in patients from the RE-VERSE AD study (N=486)

Restoration of haemostasis was achieved in 80.3% of evaluable patients who had serious bleeding and normal haemostasis was observed in 93.4% of patients who required an urgent procedure.

Of the total 503 patients, 101 patients died; each of these deaths could be attributed either as a complication of the index event or associated with co-morbidities. Thrombotic events were reported in 34 patients (23 out of the 34 patients were not on antithrombotic therapy at the time of the event) and in each of these cases, the thrombotic event could be attributed to the underlying medical condition of the patient. Mild symptoms of potential hypersensitivity (pyrexia, bronchospasm, hyperventilation, rash or pruritus) were reported. A causal relationship to idarucizumab could not be established.

5.2 Pharmacokinetic properties

The pharmacokinetics of idarucizumab were investigated in healthy subjects aged 45 to 64 years receiving a dose of 5 g as an intravenous infusion.
Distribution
Idarucizumab exhibited multiphasic disposition kinetics and limited extravascular distribution. Following the intravenous infusion of a 5 g dose, the geometric mean volume of distribution at steady state (Vss) was 8.9 L (geometric coefficient of variation (gCV) 24.8%). In the terminal phase, the volume of distribution (Vz) was 41.8 L (gCV 22.3%).

Biotransformation
Several pathways have been described that may contribute to the metabolism of antibodies. All of these pathways involve biodegradation of the antibody to smaller molecules, i.e. small peptides or amino acids which are then reabsorbed and incorporated in the general protein synthesis.

Elimination
Idarucizumab was rapidly eliminated with a total clearance of 47.0 mL/min (gCV 18.4%), an initial half-life of 47 minutes (gCV 11.4%) and a terminal half-life of 10.3 hours (gCV 18.9%). After intravenous administration of 5 g idarucizumab, 32.1% (gCV 60.0%) of the dose was recovered within a collection period of 6 hours and less than 1% in the following 18 hours. The remaining part of the dose is assumed to be eliminated via protein catabolism, mainly in the kidney.

After treatment with idarucizumab proteinuria has been observed. The transient proteinuria is a physiologic reaction to renal protein overflow after bolus/short term application of 5 g idarucizumab intravenously. The transient proteinuria usually peaked about 4 hours after idarucizumab administration and normalised within 12-24 hours. In single cases the transient proteinuria persisted for more than 24 hours.

Patients with renal impairment
In Phase I studies Praxbind has been investigated in subjects with a creatinine clearance ranging from 44 to 213 mL/min. Subjects with a creatinine clearance below 44 mL/min have not been studied in Phase I.

Depending on the degree of renal impairment the total clearance was reduced compared to healthy subjects, leading to an increased exposure of idarucizumab.

Based on pharmacokinetic data from 347 patients with different degrees of renal function (median creatinine clearance 21 - 99 mL/min) it is estimated that mean idarucizumab exposure (AUC₀–₂₄h) increases by 38% in patients with mild (CrCl 50-<80 mL/min), by 90% in moderate (30-<50 mL/min) and by 146% in severe (0-<30 mL/min) renal impairment. Since dabigatran is also excreted primarily via the kidneys, increases in the exposure to dabigatran are also seen with worsening renal function.

Based on these data and the extent of reversal of the anticoagulant effect of dabigatran in patients, renal impairment does not impact the reversal effect of idarucizumab, although the conclusion for patients with severe renal impairment is drawn from only a small subset of patients.

Patients with hepatic impairment
An impact of hepatic impairment, assessed by hepatic injury as determined by elevated liver function tests, on the pharmacokinetics of idarucizumab has not been observed. No dose adjustment is required in patients with hepatic injury.

Idarucizumab has been studied in 58 patients with varying degrees of hepatic impairment. Compared to 272 patients without hepatic impairment, the median AUC of idarucizumab was changed by -8%, 37% and 10% in patients with AST/ALT elevations of 1 to <2x ULN (N=34), 2 to <3x ULN (N=3) and >3x ULN (N=21), respectively. Based on pharmacokinetic data from 12 patients with liver disease, the AUC of idarucizumab was increased by 10% as compared to patients without liver disease.
**Older people/Sex/Race**
Based on population pharmacokinetic analyses, sex, age, and race do not have a clinically meaningful effect on the pharmacokinetics of idarucizumab.

### 5.3 Preclinical safety data

Preclinical data reveal no special hazard for humans based on repeated dose toxicity studies of up to four weeks in rats and two weeks in monkeys. Safety pharmacology studies have demonstrated no effects on the respiratory, central nervous or cardiovascular system.

Studies to evaluate the mutagenic and carcinogenic potential of idarucizumab have not been performed. Based on its mechanism of action and the characteristics of proteins no carcinogenic or genotoxic effects are anticipated.

Studies to assess the potential reproductive effects of idarucizumab have not been performed. No treatment-related effects have been identified in reproductive tissues of either sex during repeat dose intravenous toxicity studies of up to four weeks in the rat and two weeks in monkeys. Additionally, no idarucizumab binding to human reproductive tissues was observed in a tissue cross-reactivity study. Therefore, preclinical results do not suggest a risk to fertility or embryo-foetal development.

No local irritation of the blood vessel was observed after i.v. or paravenous administration of idarucizumab. The idarucizumab formulation did not produce hemolysis of human whole blood in vitro.

### 6. PHARMACEUTICAL PARTICULARS

#### 6.1 List of excipients

- glacial acetic acid
- polysorbate 20
- sodium acetate trihydrate
- sorbitol
- water for injection

#### 6.2 Incompatibilities

Praxbind must not be mixed with other medicinal products.

#### 6.3 Shelf life

36 months

Once solution has been removed from the vial, chemical and physical in-use stability of idarucizumab has been demonstrated for up to 6 hours at room temperature. The solution should not be exposed to light for more than 6 hours.

#### 6.4 Special precautions for storage

Store at 2-8°C. Do not freeze. Store in the original package in order to protect from light.

Prior to use, the unopened vial may be kept at room temperature (up to 30°C) for up to 48 hours, if stored in the original package in order to protect from light.
For storage conditions after opening of the medicinal product, see section 6.3.

6.5 Nature and contents of container

50 mL solution in a glass vial (type I glass), with a butyl rubber stopper, an aluminium cap and a label with integrated hanger.

Praxbind is supplied in packs of two vials.

6.6 Special precautions for disposal and other handling

Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration.

Praxbind must not be mixed with other medicinal products. A pre-existing intravenous line may be used for administration of Praxbind. The line must be flushed with sterile sodium chloride 9 mg/mL (0.9 %) solution prior to and at the end of infusion. No other infusion should be administered in parallel via the same intravenous access.

Praxbind is for single-use only and does not contain preservatives.

No incompatibilities between Praxbind and polyvinyl chloride, polyethylene or polyurethane infusion sets or polypropylene syringes have been observed.

7. MEDICINE SCHEDULE

Prescription Medicine

8. SPONSOR

Boehringer Ingelheim (N.Z.) Limited
P.O. Box 76-216
Manukau City
Auckland
NEW ZEALAND

Telephone: 0800 802 461

9. DATE OF FIRST APPROVAL

10 December 2015

10. DATE OF REVISION OF THE TEXT

11 June 2019

SUMMARY TABLE OF CHANGES

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<tr>
<td>4.4</td>
<td>Addition of statement regarding traceability of biological medicinal products</td>
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<tr>
<td>8</td>
<td>Sponsor contact number updated</td>
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