NEW ZEALAND DATA SHEET

1. PRODUCT NAME
ZOLADEX® 10.8 mg Depot Injection

2. QUALITATIVE AND QUANTITATIVE COMPOSITION
Goserelin (present as goserelin acetate) 10.8 mg injection.

3. PHARMACEUTICAL FORM
Injection (depot)
A sterile, white to cream coloured cylindrical depot in which goserelin acetate (equivalent to 10.8 mg of peptide base) is dispersed in a biodegradable matrix. It is supplied in a single dose syringe applicator. The SafeSystem™ incorporates a protective needle sleeve that automatically locks in place following administration of the implant to aid in the prevention of needle stick injury.

4. CLINICAL PARTICULARS

4.1 THERAPEUTIC INDICATIONS
ZOLADEX 10.8 mg is indicated for the management of:

1. Prostate cancer suitable for hormonal manipulation.

2. Adjuvant and neoadjuvant therapy in combination with radiotherapy for the management of locally advanced prostate cancer in men suitable for hormonal manipulation.

3. Endometriosis: ZOLADEX alleviates symptoms including pain, and reduces the size and number of endometrial lesions.

4. Uterine fibroids: ZOLADEX shrinks the lesions, reduces symptoms including pain, and causes cessation of menses in the majority of patients thereby improving haematological status when previous heavy menstrual loss has caused anaemia.

4.2 DOSE AND METHOD OF ADMINISTRATION
Caution should be taken while inserting ZOLADEX into the anterior abdominal wall due to the proximity of underlying inferior epigastric artery and its branches.

Use extra care when administering ZOLADEX to patients with a low BMI and/or who are receiving full anticoagulation medication (see section 4.4).
For correct administration of ZOLADEX, see instructions on the administration card (also see Method of administration below).

**Adult Men**
One depot of ZOLADEX 10.8 mg injected subcutaneously into the anterior abdominal wall every 3 months.

Adjuvant and/or neoadjuvant ZOLADEX therapy in combination with radiotherapy may include short-term use of an anti-androgen to prevent flare (see section 5.1).

**Adult Women**
One depot of ZOLADEX 10.8 mg injected subcutaneously into the anterior abdominal wall every 12 weeks.

**Children**
ZOLADEX 10.8 mg is not indicated for use in children.

**Elderly**
No dosage adjustment is necessary in the elderly.

**Renal and hepatic Impairment**
No dosage adjustment is necessary for patients with renal or hepatic impairment.

**Method of administration**
For correct administration of ZOLADEX, see instructions on the pouch/carton.

Use as directed by the prescriber. Use extra care when administering ZOLADEX to patients with a low BMI and/or who are receiving full anticoagulation medication (see section 4.4).

Use only if pouch is undamaged. Use immediately after opening pouch.

The following information is intended for medical or healthcare professionals only:

**ZOLADEX is administered by subcutaneous injection - read and understand all the instructions fully prior to administration**

1. Put the patient in a comfortable position with the upper part of the body slightly raised.
   Prepare the injection site according to the local policy and procedure.

   NOTE: Caution should be taken while injecting ZOLADEX into the anterior abdominal wall due to the proximity of underlying inferior epigastric artery and its branches; very thin patients may be at higher risk of vascular injury.

2. Examine the foil pouch and syringe for damage. Remove the syringe from the opened foil pouch and hold the syringe at a slight angle to the light.
   Check that at least part of the ZOLADEX implant is visible. (Figure 1).
3. Grasp the plastic safety tab and pull away from the syringe, and discard. (Figure 2).
Remove needle cover. Unlike liquid injections, there is no need to remove air bubbles as attempts to do so may displace the ZOLADEX implant.

4. Holding the syringe around the protective sleeve, using an aseptic technique, pinch the patient’s skin and insert the needle at a slight angle (30 to 45 degrees) to the skin.
With the opening of the needle facing up, insert needle into the subcutaneous tissue of the anterior abdominal wall below the navel line, until the protective sleeve touches the patient’s skin. (Figure 3).

NOTE: The ZOLADEX syringe cannot be used for aspiration. If the hypodermic needle penetrates a large vessel, blood will be seen instantly in the syringe chamber. If a vessel is penetrated, withdraw the needle and immediately control any resultant bleeding, monitoring the patient for signs or symptoms of abdominal haemorrhage. After ensuring the patient is haemodynamically stable another ZOLADEX implant may be injected with a new syringe elsewhere. Use extra care when administering ZOLADEX to patients with a low BMI and/or to patients receiving full dose anticoagulation.

5. Do not penetrate into muscle or peritoneum. Incorrect grip and angle of presentation is shown (Figure 4.)
6. Depress the plunger **fully**, until you can depress no more, to discharge the ZOLADEX implant and to activate the protective sleeve. You may hear a ‘click’ and will feel the protective sleeve automatically begin to slide to cover the needle. If the plunger is not depressed fully, the protective sleeve will **NOT** activate.

**NOTE**: The needle does not retract.

7. Holding the syringe as shown in **Figure 5**, withdraw the needle and allow protective sleeve to continue to slide and cover needle.
   1. Dispose of the syringe in an approved sharps collector.

![Figure 4.](image1)

NOTE: **Figure 5.**

**NOTE**: In the unlikely event of the need to surgically remove a ZOLADEX implant, it may be localized by ultrasound.

### 4.3 CONTRAINDICATIONS

Known severe hypersensitivity to the active substance or to any of the excipients of this product.

Pregnancy and lactation (see section 4.6).

### 4.4 SPECIAL WARNINGS AND PRECAUTIONS FOR USE

ZOLADEX 10.8 mg is not indicated for use in children, as safety and efficacy have not been established in this group of patients.

Injection site injury has been reported with ZOLADEX, including events of pain, haematoma, haemorrhage and vascular injury. Monitor affected patients for signs or symptoms of abdominal haemorrhage. In very rare cases, administration error resulted in vascular injury and haemorrhagic shock requiring blood transfusions and surgical intervention. Extra care
should be taken when administering ZOLADEX to patients with a low BMI and/or receiving full anticoagulation medications.

The use of ZOLADEX 10.8 mg in men at particular risk of developing ureteric obstruction or spinal cord compression should be considered carefully and the patients monitored closely during the first month of therapy. If spinal cord compression or renal impairment due to ureteric obstruction are present or develop, specific standard treatment of these complications should be instituted.

Initially ZOLADEX 10.8 mg, like other LHRH agonists, transiently increases serum testosterone. Some patients may experience a temporary increase in bone pain, which can be managed symptomatically.

A reduction in glucose tolerance has been observed in males receiving LHRH agonists. This may manifest as diabetes or loss of glycaemic control in those with pre-existing diabetes mellitus. Consideration should therefore be given to monitoring blood glucose.

An increased risk of developing myocardial infarction and, sudden cardiac death has been reported in association with use of GnRH agonists in men. The risk appears low based on the reported odds ratios, and should be evaluated carefully along with cardiovascular risk factors when determining a treatment for patients with prostate cancer. Patients receiving a GnRH agonist should be monitored for symptoms and signs suggestive of development of cardiovascular disease.

The use of LHRH agonists may cause a reduction in bone mineral density. In women, current available data suggest that recovery of bone loss occurs on cessation of therapy in the majority. In patients receiving ZOLADEX 3.6 mg for the treatment of endometriosis, the addition of hormone replacement therapy (a daily oestrogenic agent and a progestogenic agent) has been shown to reduce bone mineral loss and vasomotor symptoms. There is no experience of the use of hormone replacement therapy in women receiving ZOLADEX 10.8 mg. In men, preliminary data suggest the use of a bisphosphonate in combination with a LHRH agonist may reduce bone mineral loss.

In women, ZOLADEX 10.8 mg is only indicated for use in endometriosis and fibroids. For female patients requiring treatment with goserelin for other conditions, refer to the prescribing information for ZOLADEX 3.6 mg.

Time to return of menses after cessation of therapy with ZOLADEX 10.8 mg may be prolonged in some patients.

The use of ZOLADEX may cause an increase in cervical resistance and care should be taken when dilating the cervix.

There are no clinical data on the effects of treating benign gynaecological conditions with ZOLADEX for periods in excess of six months.

Androgen deprivation therapy may prolong the QT interval, although a causal association has not been established with ZOLADEX. In patients with a history of or who have risk factors for QT prolongation and in patients receiving concomitant medicinal products that may prolong the QT interval (see section 4.5) physicians should assess the benefit risk ratio including the potential for Torsade de Pointes prior to initiating ZOLADEX.
4.5 INTERACTION WITH OTHER MEDICINES AND OTHER FORMS OF INTERACTION

None known.

Since androgen deprivation treatment may prolong the QT interval, the concomitant use of ZOLADEX with medicinal products known to prolong the QT interval or medicinal products able to induce Torsade de Pointes should be carefully evaluated (see section 4.4).

4.6 FERTILITY, PREGNANCY AND LACTATION

Pregnancy

ZOLADEX 10.8 mg should not be used in pregnancy as there is a theoretical risk of abortion or foetal abnormality if LHRH agonists are used during pregnancy. Potentially fertile women should be examined carefully before treatment to exclude pregnancy. Non hormonal methods of contraception should be employed during therapy until menses is resumed (see section 4.4).

Breast-feeding

The use of ZOLADEX 10.8 mg during breast feeding is not recommended.

4.7 EFFECTS ON ABILITY TO DRIVE AND USE MACHINES

There is no evidence that ZOLADEX 10.8 mg results in impairment of ability to drive or operate machinery.

4.8 UNDESIRABLE EFFECTS

The following frequency categories for adverse drug reactions (ADRs) were calculated based on reports from ZOLADEX clinical trials and post-marketing sources.

<table>
<thead>
<tr>
<th>Frequency Descriptor</th>
<th>SOC</th>
<th>Males</th>
<th>Females</th>
</tr>
</thead>
<tbody>
<tr>
<td>Very Common (≥10%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Psychiatric disorders</td>
<td></td>
<td>Libido decreased⁵</td>
<td>Libido decreased⁵</td>
</tr>
<tr>
<td>Vascular disorders</td>
<td></td>
<td>Hot flush³</td>
<td>Hot flush³</td>
</tr>
<tr>
<td>Skin and subcutaneous tissue disorders</td>
<td></td>
<td>Hyperhidrosis⁵</td>
<td>Hyperhidrosis⁵, acne⁴</td>
</tr>
<tr>
<td>Reproductive system and breast disorders</td>
<td></td>
<td>Erectile dysfunction</td>
<td>N/A</td>
</tr>
<tr>
<td>Reproductive system and breast disorders</td>
<td></td>
<td>N/A</td>
<td>Vulvovaginal dryness</td>
</tr>
<tr>
<td>Reproductive system and breast disorders</td>
<td></td>
<td>N/A</td>
<td>Breast enlargement</td>
</tr>
<tr>
<td>General disorders and administration site conditions</td>
<td></td>
<td>(see Common)</td>
<td>Injection site reactions</td>
</tr>
<tr>
<td>Frequency Descriptor</td>
<td>SOC</td>
<td>Males</td>
<td>Females</td>
</tr>
<tr>
<td>----------------------</td>
<td>-----</td>
<td>-------</td>
<td>---------</td>
</tr>
<tr>
<td><strong>Common</strong> (≥1% and &lt;10%)</td>
<td>Metabolism and nutrition disorders</td>
<td>Glucose tolerance impaired(\text{\textsuperscript{b}})</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>Psychiatric disorders</td>
<td>Mood swings</td>
<td>Mood altered, depression</td>
</tr>
<tr>
<td></td>
<td>Nervous system disorders</td>
<td>Paraesthesia</td>
<td>Paraesthesia</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Spinal cord compression</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>N/A</td>
<td>Headache</td>
</tr>
<tr>
<td></td>
<td>Cardiac disorders</td>
<td>Cardiac failure(\text{\textsuperscript{f}})</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Myocardial infarction(\text{\textsuperscript{f}})</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>Vascular disorders</td>
<td>Blood pressure abnormal(\text{\textsuperscript{c}})</td>
<td>Blood pressure abnormal(\text{\textsuperscript{c}})</td>
</tr>
<tr>
<td></td>
<td>Skin and subcutaneous tissue disorders</td>
<td>Rash(\text{\textsuperscript{d}})</td>
<td>Rash(\text{\textsuperscript{d}}), alopecia(\text{\textsuperscript{g}})</td>
</tr>
<tr>
<td></td>
<td>Musculoskeletal, connective tissue and bone disorders</td>
<td>Bone pain(\text{\textsuperscript{a}})</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(see Uncommon)</td>
<td>Arthralgia</td>
</tr>
<tr>
<td></td>
<td>Reproductive system and breast disorders</td>
<td>Gynaecomastia</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>General disorders and administration site conditions</td>
<td>N/A</td>
<td>Tumour flare, tumour pain</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Injection site reaction</td>
<td>(see Very Common)</td>
</tr>
<tr>
<td><strong>Investigations</strong></td>
<td>Density decreased, weight increased</td>
<td>Bone density decreased, weight increased</td>
<td></td>
</tr>
<tr>
<td><strong>Uncommon</strong> (≥0.1% and &lt;1%)</td>
<td>Immune system disorders</td>
<td>Drug hypersensitivity</td>
<td>Drug hypersensitivity</td>
</tr>
<tr>
<td></td>
<td>Musculoskeletal, connective tissue and bone disorders</td>
<td>Arthralgia</td>
<td>(see Common)</td>
</tr>
<tr>
<td></td>
<td>Renal and urinary disorders</td>
<td>Ureteric obstruction</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>Reproductive system and breast disorders</td>
<td>Breast tenderness</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Rare</strong> (≥0.01% and &lt;0.1%)</td>
<td>Immune system disorders</td>
<td>Anaphylactic reaction</td>
<td>Anaphylactic reaction</td>
</tr>
<tr>
<td></td>
<td>Reproductive system and breast disorders</td>
<td>N/A</td>
<td>Ovarian cyst</td>
</tr>
<tr>
<td><strong>Very Rare</strong> (&lt;0.01%)</td>
<td>Neoplasms benign, malignant and unspecified (including cysts and polyps)</td>
<td>Pituitary tumour</td>
<td>Pituitary tumour</td>
</tr>
<tr>
<td></td>
<td>Endocrine disorders</td>
<td>Pituitary haemorrhage</td>
<td>Pituitary haemorrhage</td>
</tr>
</tbody>
</table>
Frequency Descriptor | SOC | Males | Females |
--- | --- | --- | --- |
Psychiatric disorders | Psychotic disorder | Psychotic disorder |
Unknown | Neoplasms benign, malignant and unspecified (including cysts and polyps) | N/A | Degeneration of uterine fibroid |
Skin and subcutaneous tissue disorders | Alopecia<sup>b</sup> | (see Common) |

<sup>a</sup> These are pharmacological effects which seldom require withdrawal of therapy.

<sup>b</sup> A reduction in glucose tolerance has been observed in males receiving LHRH agonists. This may manifest as diabetes or loss of glycaemic control in those with pre-existing diabetes mellitus.

<sup>c</sup> These may manifest as hypotension or hypertension, have been occasionally observed in patients administered ZOLADEX. The changes are usually transient, resolving either during continued therapy or after cessation of therapy with ZOLADEX. Rarely, such changes have been sufficient to require medical intervention, including withdrawal of treatment from ZOLADEX.

<sup>d</sup> These are generally mild, often regressing without discontinuation of therapy.

<sup>e</sup> Initially, prostate cancer patients may experience a temporary increase in bone pain, which can be managed symptomatically.

<sup>f</sup> Observed in a pharmaco-epidemiology study of LHRH agonists used in the treatment of prostate cancer. The risk appears to be increased when used in combination with anti-androgens.

<sup>g</sup> Loss of head hair has been reported in females, including younger patients treated for benign conditions. This is usually mild but occasionally can be severe.

<sup>h</sup> Particularly loss of body hair, an expected effect of lowered androgen levels.

<sup>i</sup> In most cases acne was reported within one month after the start of ZOLADEX.

Reduction in glucose tolerance, manifesting as diabetes or loss of glycaemic control in those with pre-existing diabetes, has been reported during treatment with GnRH agonists including ZOLADEX (see section 4.4).

A small increased risk of developing myocardial infarction and, sudden cardiac death has been reported in association with use of GnRH agonists in men.

**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 limited experience of overdosage in humans. In cases where ZOLADEX has unintentionally been re-administered early or given at a higher dose, no clinically relevant adverse effects have been seen. Animal tests suggest that no effect other than the intended therapeutic effects on sex hormone concentrations and on the reproductive tract will be evident with higher doses of ZOLADEX 10.8 mg. If overdosage occurs, this should be managed symptomatically.

For advice on the management of overdose please contact the National Poisons Centre on 0800 POISON (0800 764 766).
5. PHARMACOLOGICAL PROPERTIES

5.1 PHARMACODYNAMIC PROPERTIES

ZOLADEX (D-Ser(But)⁶Azgly⁴ LHRH) is a synthetic analogue of naturally occurring luteinising-hormone releasing hormone (LHRH). On chronic administration, ZOLADEX 10.8 mg results in inhibition of pituitary luteinising-hormone secretion leading to a fall in serum testosterone concentrations in men and serum oestradiol concentrations in women. Initially ZOLADEX 10.8 mg, like other LHRH agonists, may transiently increase serum testosterone concentrations in men and serum oestradiol concentrations in women.

In men by around 21 days after the first depot injection, testosterone concentrations have fallen to within the castrate range and remain suppressed with treatment every 3 months. If in exceptional circumstances repeat dosing does not occur at 3 months, data indicate that castrate levels of testosterone are maintained for up to 16 weeks in the majority of patients.

In women, serum oestradiol concentrations are suppressed by around 4 weeks after the first depot injection and remain suppressed until the end of the treatment period. In patients with oestradiol already suppressed by an LHRH analogue, suppression is maintained on the change of therapy to ZOLADEX 10.8 mg. Suppression of oestradiol is associated with a response in endometriosis and uterine fibroids and will result in amenorrhoea in the majority of patients.

During early treatment with ZOLADEX some women may experience vaginal bleeding of variable duration and intensity. Such bleeding probably represents oestrogen withdrawal bleeding and is expected to stop spontaneously.

During treatment with LHRH analogues, patients may enter the natural menopause. Rarely, some women do not resume menses on cessation of therapy.

Clinical efficacy and safety

Effect in prostate cancer – Adjuvant and neoadjuvant ZOLADEX therapy in combination with radiotherapy

Four phase III, open-labelled, randomised, controlled, multi-centred clinical trials have been conducted to evaluate the added value of adjuvant and/or neoadjuvant ZOLADEX therapy in combination with radiotherapy in patients with histologically proven prostate cancer. The majority of patients had locally advanced disease (T2 N+, T3 or T4, N0/Nx, M0). All studies have been performed by two independent collaborative oncology groups (European Organisation for Research and Treatment of Cancer [EORTC] and the Radiation Therapy Oncology Group [RTOG]), and have reported results from median follow-up of more than 5 years. Table 2 summarises the study design, patient populations and median follow-up periods for these studies.
Table 2: Study design, patient population and median follow-up period for adjuvant and/or neoadjuvant ZOLADEX combined with radiotherapy clinical trials.

<table>
<thead>
<tr>
<th>Trial</th>
<th>Adjuvant</th>
<th>Neo-adjuvant</th>
<th>Neo and adjuvant</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>RTOG 85-31 (n=945)</td>
<td>EORTC 22863 (n=415)</td>
<td>RTOG 86-10 (n=456)</td>
</tr>
</tbody>
</table>
| Treatment     | ZOLADEX* + RT             | ZOLADEX** + RT             | ZOLADEX** + RT              | ZOLADEX** (neo) + RT + ZOLADEX*
|               |                           |                            | alone (adjuvant)            |
| Comparator    | RT alone + ZOLADEX at relapse | RT alone                  | RT alone                   | ZOLADEX** (neo) only + RT    |
| Duration      | Last week of RT continued indefinitely | Day 1 of RT continued for 3 years post RT | 2 months prior to & during RT | 2 months prior to, during & 2 years post RT |
|               |                           |                            |                            | 2 months prior to & during RT |
| Patient population | T1-2N+ & T3 (any N); Lesions <25cm³; prior prostatectomy allowed^ | T1-2N0-X (G3) & T3-4N0 (any G) | T2b-4M0; N+ allowed†; Lesions ≥25cm³ | T2c-T4; PSA <150ng/mL; N+ allowed†; KS ≥70 |
| Median follow-up | 7.6 years^a              | 5.5 years^b                | 6.7 years^c                | 5.8 years^d                 |

T, N – Tumour, node in accordance with the UICC classification; G – WHO grade; *3.6 mg sc every 4 weeks; # plus 1 month of oral cyproterone acetate 150mg/day initiated 1 week prior to Zoladex to prevent flare; RT – radiotherapy; § combined with oral flutamide (250mg three times daily); ^ if penetration to the margins of resection and/or seminal vesicle involvement + Karnofsky performance status >60%; † if below the common iliac chain; KS – Karnofsky score.


Adjuvant ZOLADEX therapy long term (≥3 years) significantly improved disease-free survival and overall survival compared to radiotherapy alone (Tables 3 and 4). Neoadjuvant ZOLADEX therapy for two months prior and during radiotherapy significantly improved disease-free survival but not overall survival compared to radiotherapy alone (Table 5). A combination of neoadjuvant and adjuvant therapy (2 years) also significantly improved disease-free survival but not overall survival compared to radiotherapy alone (Table 6).

Table 3: Adjuvant ZOLADEX efficacy results for RTOG 85-31 (median follow-up: all patients 7.6 years; alive patients 10 years)

<table>
<thead>
<tr>
<th>Endpoint</th>
<th>10 year estimates (%)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ZOLADEX + RT</td>
<td>RT alone</td>
</tr>
<tr>
<td>Overall survival</td>
<td>47*</td>
<td>38</td>
</tr>
<tr>
<td>Disease-free survival</td>
<td>30</td>
<td>9</td>
</tr>
</tbody>
</table>

*ASCO presentation slides
Table 4: Adjuvant ZOLADEX efficacy results for EORTC 22863 (median follow-up: all patients 5.5 years)

<table>
<thead>
<tr>
<th>Endpoint</th>
<th>5 year estimates (%)</th>
<th>Hazard ratio [95% CI]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ZOLADEX +RT</td>
<td>RT alone</td>
</tr>
<tr>
<td>Overall survival</td>
<td>78</td>
<td>62</td>
</tr>
<tr>
<td>Disease-free survival</td>
<td>74</td>
<td>40</td>
</tr>
</tbody>
</table>

CI – confidence interval

Table 5: Neoadjuvant ZOLADEX efficacy results for RTOG 86-10 (median follow-up: all patients 6.7 years; alive patients 8.6 years)

<table>
<thead>
<tr>
<th>Endpoint</th>
<th>8 year estimates (%)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ZOLADEX + RT</td>
<td>RT alone</td>
</tr>
<tr>
<td>Overall survival</td>
<td>53 {53*}</td>
<td>44 {43*}</td>
</tr>
<tr>
<td>Disease-free survival</td>
<td>49</td>
<td>34</td>
</tr>
</tbody>
</table>

*updated analyses (Shipley et al 2002 – all patients 6.7 years; alive patients 9.0 years)

Table 6: Neoadjuvant and/or adjuvant ZOLADEX efficacy results for the total RTOG 92-02 population (median follow-up: all patients 5.8 years; alive patients 6.3 years)

<table>
<thead>
<tr>
<th>Endpoint</th>
<th>5 year estimates (%)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Neo &amp; adjuvant ZOLADEX</td>
<td>Neo ZOLADEX only</td>
</tr>
<tr>
<td>Overall survival</td>
<td>80.0</td>
<td>78.5</td>
</tr>
<tr>
<td>Disease-free survival</td>
<td>46</td>
<td>28</td>
</tr>
</tbody>
</table>

ns – not significant

5.2 PHARMACOKINETIC PROPERTIES

Administration of ZOLADEX 10.8 mg in accordance with the dosage recommendations ensures that exposure to goserelin is maintained with no clinically significant accumulation. ZOLADEX is poorly protein bound and has a serum elimination half-life of 2 to 4 hours in subjects with normal renal function. The half-life is increased in patients with impaired renal function. For the compound given, as recommended in a 10.8 mg depot formulation, this change will not lead to any accumulation. Hence, no change in dosing is necessary in these patients. There is no significant change in pharmacokinetics in patients with hepatic failure.

5.3 PRECLINICAL SAFETY DATA

Following long-term repeated dosing with ZOLADEX, an increased incidence of benign pituitary tumours has been observed in male rats. While this finding is similar to that previously noted in this species following surgical castration, any relevance to humans has not been established.

In mice, long term repeated dosing with multiples of the human dose produced histological changes in some regions of the digestive system manifested by pancreatic islet cell
hyperplasia and a benign proliferative condition in the pyloric region of the stomach, also reported as a spontaneous lesion in this species. The clinical relevance of these findings is unknown.

ZOLADEX is a synthetically derived peptide.

6. PHARMACEUTICAL PARTICULARS

6.1 LIST OF EXCIPIENTS
Polyglactin

6.2 INCOMPATIBILITIES
Not applicable.

6.3 SHELF LIFE
3 years

6.4 SPECIAL PRECAUTIONS FOR STORAGE
Store below 25°C.

6.5 NATURE AND CONTENTS OF CONTAINER
There is one depot per pack.

6.6 SPECIAL PRECAUTIONS FOR DISPOSAL
Return unused and expired medicines to your local pharmacy for disposal.

7. MEDICINE SCHEDULE
Prescription Medicine.

8. SPONSOR
AstraZeneca Limited
PO Box 87453
Meadowbank
Auckland 1742.
Telephone: (09) 306 5650

9. DATE OF FIRST APPROVAL
12 September 1996
10. DATE OF REVISION OF THE TEXT
8 May 2020
CDS 130515+ Aust PI

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SUMMARY TABLE OF CHANGES

<table>
<thead>
<tr>
<th>Section changed</th>
<th>Summary of new information</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>PO Box address updated.</td>
</tr>
<tr>
<td>10.</td>
<td>Trademark information updated.</td>
</tr>
</tbody>
</table>