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- 4
- 5 Title
- 6 Denosumab for treatment of bone metastases secondary to solid tumours:
- 7 systematic review and network meta-analysis
- 8

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# 29 Abstract

- 30 Aim
- 31 To evaluate the evidence for denosumab for the treatment of bone metastases secondary to
- 32 solid tumours and, using a network meta-analysis, indirectly compare denosumab with
- 33 bisphosphonates and best supportive care.
- 34 Data sources
- 35 MEDLINE (1948 to April 2011), EMBASE (1980 to March 2011), Cochrane Library (all
- 36 sections) (Issue 1, 2011) and Web of Science with Conference Proceedings (1970 to May
- 2011) and additional meeting abstracts (2010 and 2011) were searched.
- 38 Study eligibility, participants and interventions
- 39 Only randomised controlled trials assessing denosumab, bisphosphonates or best
- 40 supportive care in patients with bone metastases from any solid tumour were included.
- 41 Synthesis
- 42 Direct evidence comparing denosumab and zoledronic acid was assessed for breast cancer,
- 43 prostate cancer and other solid tumours. Denosumab was compared with pamidronate and
- 44 best supportive care through a network meta-analysis for each tumour type. The primary
- 45 outcomes were time to first skeletal related event (SRE) and time to first and subsequent
- 46 SRE. Secondary outcomes were skeletal morbidity rate, pain, quality of life (QoL) and overall
- 47 survival.
- 48 Results
- 49 Denosumab was found to be more effective in delaying the time to first SRE and reducing
- 50 the risk of first and subsequent SREs compared to zoledronic acid, placebo and
- 51 pamidronate. In breast and prostate cancer, denosumab was effective in reducing skeletal
- 52 morbidity rate compared with placebo. The lack of published data on pain and QoL meant
- that firm conclusions could not be made. Denosumab did not appear to have an affect on
- 54 overall survival.
- 55 Limitations
- 56 Network meta-analyses are subject to uncertainties and potential biases.
- 57 Conclusions
- 58 Denosumab is effective in preventing SREs, but the effect on pain and QoL is unclear.

# 59 Key words

- 60 upto 10 MESH keywords
- 61 denosumab, zoledronic acid, pamidronate, neoplasm metastasis, indirect estimation
- 62 techniques

#### 63 Introduction

- 64 The impact of bone metastases on cancer patients can be considerable. Complications,
- reduced mobility, pain and the effects of treatment reduce quality of life significantly.
- 66 Complications may include pathological fracture, spinal cord compression and
- 67 hypercalcaemia of malignancy.

68 Bone-targeted pharmacological treatments aim at preventing complications, reducing pain

- and improving quality of life. To date bisphosphonates have been the main pharmacological
- 70 treatment option for patients with bone metastases. Currently licensed bisphosphonates
- include; zoledronic acid (any advanced malignancy involving bone), disodium pamidronate
- 72 (breast cancer or multiple myeloma), sodium clodronate (breast cancer or multiple myeloma)
- and ibandronic acid (breast cancer). Bisphosphonates are administered either intravenously
- 74 (zoledronic acid, pamidronate or ibandronic acid) or orally (clodronate or ibandronic acid)
- and have been associated with renal toxicity.<sup>1</sup> In the UK, the National Institute of Health and
- 76 Clinical Excellence (NICE) currently recommends the use of bisphosphonates in all patients
- with bone metastases secondary to breast cancer,<sup>2</sup> patients with hormone resistant prostate
- cancer with painful bone metastases despite conventional analgesics<sup>3</sup> or as an option in
- <sup>79</sup> lung cancer with bone metastases.<sup>4</sup> Patients who are not recommended for
- 80 bisphosphonates would receive standard best supportive care.
- 81 Denosumab (Xgeva, Amgen) is a fully human monoclonal antibody, licensed for the
- 82 prevention of skeletal related events (SRE) in bone metastases from solid tumours. It is
- 83 administered by sub-cutaneous injection and does not require renal monitoring.<sup>5</sup>
- The term 'skeletal related event' is a composite endpoint that has evolved over the past 20 years for use in clinical trials. Recent trials define SREs as pathological fracture (including asymptomatic vertebral collapse), spinal cord compression or need for radiotherapy or surgery to bone.<sup>6-8</sup> Other definitions have included hypercalcaemia or change in antineoplastic therapy.
- Three pivotal trials have evaluated denosumab compared to zoledronic acid for the prevention of SREs.<sup>6-8</sup> There are no head-to-head trials of denosumab compared with other bisphosphonates or best supportive care. These comparisons are, nonetheless, important because of the wide variation in practice. Some centres use only zoledronic acid, some use a variety of bisphosphonates, while others do not use bisphosphonates at all (especially in cancer other than breast). Therefore the aim of this review is to evaluate the evidence for denosumab for the treatment of bone metastases in solid tumours and, using a network

- 96 meta-analysis, indirectly compare denosumab with other bisphosphonates and best
- 97 supportive care.

#### 98 Materials and methods

99

The review complies with PRIMSA guidelines.<sup>9</sup> A pre-specified protocol has been published 100 on the NICE website.<sup>10</sup> 101

102

#### 103 Literature search and eligibility criteria

Studies were identified by systematic searching of the following databases; MEDLINE (1948 104 to April 2011), EMBASE (1980 to March 2011), Cochrane Library (all sections) (Issue 1, 105 2011) and Web of Science with Conference Proceedings (1970 to May 2011). Additional 106 107 meeting abstracts (2010 and 2011) were identified through searching American Society of Clinical Oncology, American Urological Association and San Antonio Breast Cancer 108 109 symposium. Reference lists of all included studies were scanned to identify additional potentially relevant studies. The titles and abstracts of all papers identified by the search 110 111 strategy were screened and full-text copies of all potentially relevant studies obtained. 112

113 The search strategy used for MEDLINE was; step 1) exp Diphosphonates, step 2) RANK

114 Ligand, step 3) (denosumab or bisphosphonate\* or ibandron\* or clodron\* or pamidron\* or

zoledron\*).tw., step 4) (radiation or radiotherapy or radionuclide\* or hormone therapy or 115

strontium or samarium).ti., step 5) or/1-4, step 6) exp Neoplasms, step 7) (solid tumor or 116

solid tumour\* or cancer or carcinoma or myeloma).tw., step 8) or/6-7, step 9) 5 and 8, step 117

10) exp Bone Neoplasms, step 11) (((bone or osteolytic or lytic) adj lesion\*) or (bone adj2 118

metast\*)).tw., step 12) (skeletal or fracture\*).tw., step 13) or/10-12, step 14) 9 and 13, step 119

15) randomized controlled trial.pt., step 16) 14 and 15 and, step 17) limit 16 to english 120

121 language.

122

This search strategy was adapted as appropriate for the other databases 123

124

125 Only randomised controlled trials evaluating denosumab, bisphosphonates or best supportive care were included. Best supportive care included trials evaluating radiotherapy, 126 radionuclides, hormone therapy, strontium or samarium. Bone metastases secondary to any 127 128 solid tumour were eligible.

129

130 Screening was performed by two independent authors and disagreements resolved by 131 discussion. After piloting a data extraction form, data were extracted by one author and

- 132 checked by a second. Data included study characteristics, inclusion/exclusion criteria,
- 133 results and adverse events. Quality was assessed using the Cochrane risk of bias tool.<sup>11</sup>
- 134
- 135 The primary outcomes were time to first SRE and time to first and subsequent SRE.
- 136 Secondary outcomes were skeletal morbidity rate (SMR, , defined as ratio of the number of
- 137 SREs per patient divided by the patient's time at risk), pain, quality of life and overall
- 138 survival.
- 139

#### 140 Network meta-analysis

141 Network meta-analysis (NMA) is a statistical technique used to indirectly compare two or

142 more interventions. Generally, it is used in situations where there is an absence of head-to-

143 head trials.

144 Studies meeting the inclusion criteria were assessed for eligibility of synthesis by network

145 meta-analysis, by evaluating methodological heterogeneity. To be suitable for NMA, studies

146 were required to be similar with respect to population, intervention, comparators, outcomes,

- 147 SRE definition and time frame. Based on this assessment, networks were designed.
- 148 Networks were created for three primary cancer types; breast cancer, prostate cancer and
  149 other solid tumours including (OST). A subgroup of patients with non small cell lung cancer
  150 within OST was also explored.

The analyses followed methods for mixed treatment comparisons described by Lu and
Ades.<sup>12</sup> and used the Bayesian software package, WinBUGS, which employs Markov chain
Monte Carlo (MCMC) methods.

154 Outcomes analysed were time to first SRE (hazard ratios), time to first and subsequent SRE 155 (rate ratios from Andersen-Gill<sup>13</sup> multiple event analyses reported in primary studies) and 156 SMR ratios (for breast and prostate cancer only).

Fixed effects models were used for time to first SRE, adopting an approach recommended 157 by the NICE Decision Support Unit<sup>14</sup> for modelling trial-based summary measures, which 158 can be applied to modelling hazard ratios on the log hazard scale. The trial-level data 159 included in the models comprised log hazard ratios and its standard error. Where hazard 160 ratios were not reported or derivable in the primary study or related publications (e.g. 161 publically available FDA documentation), Kaplan-Meier estimates and numbers at risk (if 162 available) were used, applying the methods of Tierney<sup>15</sup> to estimate the hazard ratio. 163 Pairwise hazard ratios were estimated from the median of the posterior distribution with 164

credible intervals taken from the 2.5% and 97.5% percentiles. Ten thousand MCMC
simulations were used in the analysis following a burn-in of 10,000. The same approach
was taken for modelling rate ratios in the analysis of time to first and subsequent SREs.

For SMR a random effects model was adopted using arm-based data. The data included in 168 169 the SMR models were mean SMR and standard deviation along with the number of patients. Where standard deviations were not reported, values were imputed by taking the mean of 170 171 reported SDs from other studies but for the same treatment. The robustness of the 172 imputation was tested by comparing results with those obtained by treating missing data as an uncertain parameter. Posterior distributions for relative treatment effects were estimated 173 174 from the absolute risks of outcome from the relevant individual treatments. Median 175 estimates and credible intervals were taken from 10,000 MCMC simulations after a burn-in of 10,000. 176

177 In order to estimate the absolute risk of outcome in the analyses of arm-based data, it was 178 necessary to include an estimate of the baseline risk of the control treatment in the models. Zoledronic acid was treated as the reference treatment in each analysis as it is the treatment 179 common to the largest number of trials and is present in multiple included studies for each 180 NMA. Single-arm meta-analyses of zoledronic acid were conducted to estimate baseline 181 182 risk from studies included in the NMA that had zoledronic acid as one of its comparators. The data in the time-to-event analyses, however, were trial-based and baseline risk could 183 not be estimated so the absolute effect of the reference treatment was set to zero in these 184 models. 185

The quality of the models was examined by inspecting convergence using Gelman-Rubin-Brooks plots, assessing autocorrelation between iterations of the Markov chain and checking whether the MC error was less than 5% of the posterior standard deviation. 189 Results

190

#### 191 Literature search

Results of the literature search are shown in figure 1. Thirty-eight studies met the inclusion
criteria, most of which compared bisphosphonates with placebo. Of these 38 studies, 30
were excluded because they were not suitable for network meta-analysis (table 1). The
characteristics and results of the eight studies included in the NMA are shown in table 2 and
3.

197

## 198 Study quality

The quality of the studies included in the NMA was high as shown in table 4. There was a
low risk of bias for the majority of categories. Stopeck 2010<sup>8</sup> and Rosen 2003<sup>16</sup> failed to
describe sequence generation or allocation concealment. Kohno 2005<sup>17</sup> and Rosen 2003<sup>16</sup>
did not sufficiently address incomplete outcome data.

203

### 204 Study characteristics

Four studies included patients with breast cancer,<sup>8,16-18</sup> two with prostate cancer<sup>6,19</sup> and two with other solid tumours<sup>7,20</sup> (table 2). Henry 2011 included patients with multiple myeloma, in addition to patients with other solid tumours. Three studies compared denosumab with zoledronic acid,<sup>6-8</sup> three compared zoledronic acid with placebo,<sup>17,19,20</sup> one zoledronic acid with pamidronate<sup>16</sup> and one pamidronate with placebo.<sup>18</sup>

Six studies were international, one study only recruited patients from Japan<sup>17</sup> and one study recruited patients from the US.<sup>18</sup> Patients were youngest in the breast cancer studies and oldest in the prostate. The proportion of patients with a previous SRE at baseline ranged from 24%<sup>6</sup> to 73%.<sup>20</sup>

214

## 215 Direct SRE results

216 Denosumab statistically significantly delayed the time to first on-study SRE in breast cancer,

217 prostate cancer and other solid tumours (table 3). The difference in mean months of time to

first SRE between denosumab and zoledronic acid was 3.6 months in prostate cancer (HR

- 219 0.82 95%CI 0.71 to 0.95) and 4.3 months in other solid tumours (HR 0.84 95%CI 0.71 to
- 220 0.98) (in breast cancer this outcome was not reached (HR 0.82 9%%CI 0.71 to 0.95)).
- 221 Similarly, denosumab statistically significantly reduced the risk of time to first and
- subsequent SRE for prostate cancer (rate ratio 0.82 95%CI 0.71 to 0.94) and breast cancer
- (rate ratio 0.77, 95%CI 0.66 to 0.89). In other solid tumours, the result favoured denosumab
- but was not statistically significant (rate ratio 0.90 95%Cl 0.77 to 1.04).
- Stopeck 2010<sup>8</sup> was the only trial evaluating denosumab to report SMR. Denosumab was
  associated with a lower SMR compared with zoledronic acid (0.45 compared with 0.58, p
  value 0.004) in patients with breast cancer.
- In the bisphosphonate trials, zoledronic acid and pamidronate were associated with delayed
- time to first SRE, time to first and subsequent SRE and SMR. In the only trial comparing
- 230 zoledronic acid and pamidronate,<sup>16</sup> the authors found that zoledronic acid statistically
- significantly reduced the time to first SRE in hormone-treated breast cancer patients (415
- days versus 370 days, p = 0.047) and risk of time to first and subsequent SRE in all breast
- 233 cancer patients (RR = 0.80 (0.66 to 0.97).
- 234

# 235 Pain study results

- 236 Stopeck 2010<sup>8</sup> reported that the median time to developing moderate/severe pain in women
- with breast cancer, in patients with no/mild pain at baseline, was longer in denosumab
- compared with zoledronic acid (295 days versus 176 days; HR 0.78, 95%Cl 0.67 to 0.92)
- 239 Pain outcomes for denosumab compared with zoledronic acid in other solid tumours is
- available in abstract form.<sup>21</sup> Denosumab was found to delay the time to clinically significant
- pain (more than 2 point increase from baseline on brief pain inventory) compared to
- zoledronic acid (169 days compared with 143 days HR 0.85, 95% CI: 0.73-0.98).
- In prostate cancer, pain data have also been published in abstract form.<sup>22</sup> In the subgroup of
  patients with no/mild pain at baseline, there was no statistically significant difference in the
  time to moderate/severe in denosumab compared to zoledronic acid (177 days versus 148
  days; HR 0.89, 95% CI 0.77, 1.04).

#### 247 Quality of life study results

<sup>248</sup> In breast cancer, quality of life data for denosumab have been published in abstract form.<sup>23</sup>

The authors report that over the 18 month period an average of 4.1% more (range -0.6% to

9.3%) patients treated with denosumab, compared with zoledronic acid, experienced a

251 meaningful improvement in quality of life (5 or more increase in FACT-G score).

No quality of life data are available for prostate cancer or other solid tumours.

253

# 254 Overall survival study results

There was no significant difference in overall survival between denosumab and zoledronic
acid in breast cancer and prostate cancer. Henry 2010<sup>24</sup> also reported no significant
difference; however on ad hoc analysis the authors found that denosumab was associated
with an increased overall survival in non small cell lung cancer (HR 0.79, 95%CI 0.65 to
0.95). Notably the authors also reported a decrease in overall survival in the ad hoc analysis
of multiple myeloma patients (HR 2.26, 95%CI 1.13 to 4.50).

261

# 262 Safety

263 For breast, prostate and other solid tumours denosumab, compared with zoledronic acid,

was associated with lower renal impairment (0.4% versus 2.2%, 16% versus 15%, 8.3%

versus 10.9%) and acute phase reaction (10.4% versus 27.3%, 8% versus 18%, 6.9%

versus 14.5%). However, denosumab was associated with higher incidence of

hypocalcaemia (not reported, 13% versus 6%, 2.3% versus 1.0%) and osteonecrosis of the

268 jaw (2.0% versus 1.4%, 2 versus 1%, 1.1% versus 1.3%).

269

#### 270 Network meta-analysis results

Network diagrams for breast cancer, prostate cancer and other solid tumours are shown in
figures 2, 3 and 4. The same network was used for the subgroup of non small cell lung
cancer as other solid tumours. The results of these analyses are summarised in tables 5, 6
and 7.

#### 275 Denosumab versus placebo

- 276 NMA results suggest that denosumab, compared with placebo, reduces the time to first SRE
- in breast, prostate cancer and other solid tumours. In non small cell lung cancer the result
- favoured denosumab, but was not statistically significant (HR 0.68, 95%CI 0.45 to 1.03).
- 279 Similarly denosumab statistically significantly reduced the risk of first and subsequent SRE in
- 280 breast cancer, prostate cancer, other solid tumours and non small cell lung cancer,
- 281 compared to placebo. Additionally, denosumab reduced the skeletal morbidity rate
- compared with placebo in all groups.

### 283 Denosumab versus pamidronate

- 284 The comparison of denosumab versus pamidronate was only possible in breast cancer. For
- skeletal morbidity rate the result favours denosumab, but there was no significant difference.
- 286 There was a significant difference in time to first SRE and time to first and subsequent SRE
- when denosumab was compared with pamidronate (HR 0.73 95%Cl 0.56 to 0.94 and rate
- 288 ratio 0.62 95%CI 0.48 to 0.80, respectively).

#### 289 Discussion

290

#### 291 Statement of key findings

Based on the review of direct evidence and network meta-analysis, denosumab, compared with zoledronic acid or placebo, statistically significantly delays time to first SRE, time to first and subsequent SRE and skeletal morbidity rate. Denosumab appears to be more effective than pamidronate for these outcomes, but the results have mixed statistical significance.

Although denosumab has demonstrated its effectiveness in delaying SREs, a lack of published data means that conclusions about pain and quality of life cannot be made. There was no statistically significant difference in overall survival for denosumab compared with zoledronic acid for prostate and breast cancer. However in an ad hoc analysis of the trial including various tumour types, denosumab was found to improve the overall survival in nonsmall cell lung cancer.

302

#### 303 Strengths and limitations

There are a number of strengths of this review. A comprehensive and robust search strategy was used. A rigorous inclusion/exclusion criteria was used which only included high quality evidence (RCTs). Undertaking a NMA means that estimates of effectiveness can be made when no direct evidence is available. This was the case for comparing denosumab with placebo and pamidronate. Excluding studies with a different definition of what constitutes an SRE resulted in a smaller but more robust NMA.

Although NMA allows indirect estimates to be calculated, they can be subject to potential 310 biases and uncertainties.<sup>25</sup> Network meta-analyses are not randomised comparisons, but 311 312 rather observational findings across studies and therefore should be interpreted with due caution. The quality of any NMA is only as good as the weakest link in the network. All 313 studies included in this NMA were of good quality (table 4), improving the validity of the NMA 314 results. Some published studies did not report full results, therefore some treatment effects 315 were estimated, for example using the method described by Teirney and colleagues.<sup>15</sup> 316 However when these parameters were treated as uncertain, the impact on the results was 317 318 negligible. A key limitation was the small number of studies included. This resulted in an 319 unstable model when a random effects model was used for time to first SRE and time to first 320 and subsequent SRE. Therefore a fixed effects model was used, which assumes no variability between studies. 321

323

#### 324 Meaning of the results

Our analysis indicates that denosumab is effective in delaying first and first-and-subsequent SREs when compared to zoledronic acid, placebo and pamidronate. NMA analysis results in reduced power and therefore less precision. Non-statistically significant results for skeletal morbidity rate for denosumab compared with pamidronate should not be interpreted as evidence that there is no effect. Only if higher powered NMA were possible could this conclusion be made.

331 The validity of these results relies on, firstly, the SRE outcome and, secondly, the analysis of 332 it. The SRE outcome is useful because it allows for increased power and therefore 333 efficiency. It would be impractical to power trials to detect differences in each component of the SRE outcome, especially with regard to spinal cord compression and need for surgery to 334 335 bone (as these are rare events). However, the composite outcome is of little use to patients since it incorporates a wide spectrum of clinical events, ranging from asymptomatic 336 pathological fracture (identified during routine on-study skeletal surveys) to paraplegic spinal 337 cord compression. Furthermore, the outcome does not directly measure mobility or bone 338 pain, although it could be argued that the need for radiotherapy is an indirect measure of 339 bone pain. In addition, for many patients, radiotherapy will be a highly effective treatment for 340 bone pain. 341

Using time to event and multiple event analyses (time to first and subsequent SRE) allows smaller differences between treatments to be identified. This may be warranted when comparing active comparators; however, researchers and healthcare staff should ensure that statistically significant differences are clinically meaningful. In addition, the method used in these trials for the multiple event analysis (Andersen-Gill<sup>13</sup>) has been criticised because it does not differentiate between participants who died and who leave the study for another reason.<sup>26</sup> These issues have been discussed in greater detail elsewhere.<sup>27</sup>

A key issue is whether the delay in SREs results in a reduction in pain and improvement in quality of life. Ideally, the improved SRE outcomes with denosumab, would be interpreted alongside pain and quality of life data. Unfortunately, the lack of published pain and quality of life data means that this association could not be established. The data published from the three pivotal trials are only available in abstract form and generally only reports subgroups. For breast cancer there was a statistically significant delay to moderate/severe pain in

322

patients with no/mild pain, however in prostate cancer the difference was not statisticallysignificant.

357 Denosumab has the added advantage of being given as a sub-cutaneous injection which 358 does not require renal monitoring. Denosumab could potentially be administered in the 359 community. Zoledronic acid is an intra-venous administration and requires renal monitoring with dose adjustment if renal impairment present. In terms of adverse events, denosumab 360 361 has lower renal toxicity and does not appear to be associated with acute phase reactions. 362 However, there is a marginally higher incidence of osteonecrosis of the jaw. In addition, there is a higher incidence of hypocalcaemia but this can be easily corrected with 363 364 appropriate treatment.

365

#### 366 Future research needs

367 In common with most findings for bisphosphonates in advanced cancer, from available

368 evidence denosumab does not appear to affect overall survival. In the Henry 2010 trial,<sup>24</sup>

369 there was a statistically significant improvement in overall survival in the ad hoc analysis for

non small cell lung cancer. The reason for this is not clear and it may be a chance finding.

371 Further trials in this subgroup would be needed to establish the validity of this result..

The place for denosumab in treatment pathways is unclear. Much of this will depend on local budgets and on economic evaluations.<sup>28,29</sup> One option may be as a second line agent in patients who suffer an SRE on bisphosphonates. A randomised controlled trial looking at this specific population may be informative.

376

# 377 Conclusion

378 Denosumab compared with zoledronic acid, placebo and, pamidronate, is effective in

delaying time to first SRE and reducing the risk of first and subsequent SRE. However,

380 conclusion about its impact on pain reduction and quality of life cannot be reached because

381 of the lack of published data.

382

# 383 Financial and disclosure statement

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## 397 References

- Ross JR, Saunders Y, Edmonds PM, Patel S, Wonderling D, Normand C et al. A
   systematic review of the role of bisphosphonates in metastatic disease. *Health Technol Assess* 2004;8:1-176.
- 401 2 CG81: Advanced breast cancer: diagnosis and treatment [document on the Internet].
  402 London: National Institute for Health and Clinical Excellence; 2009 [accessed October
  403 2011]. Available from URL:
- 404 <u>http://www.nice.org.uk/nicemedia/live/11778/43414/43414.pdf</u>.
- 405 3 CG58: Prostate cancer: diagnosis and treatment [document on the Internet]. London:
  406 National Institute for Health and Clinical Excellence; 2008 [accessed October 2011].
  407 Available from URL: <u>http://www.nice.org.uk/nicemedia/live/11924/39687/39687.pdf</u>.
- 408 4 CG121: The diagnosis and treatment of lung cancer (update) [document on the lnternet]. London: National Institute for Health and Clinical Excellence; 2011 [accessed
  410 October 2011]. Available from URL: http://www.nice.org.uk/nicemedia/live/13465/54199/54199.pdf.
- 5 Xgeva: EPAR product information. Annex 1: Summary of product characteristics [document on the Internet]. London: European Medicines Agency; 2011 [accessed April 2012]. Available from URL:
  <u>http://www.ema.europa.eu/docs/en\_GB/document\_library/EPAR\_-</u>
  <u>Product\_Information/human/002173/WC500110381.pdf</u>.
- Fizazi K, Carducci M, Smith M, Damiao R, Brown J, Karsh L et al. Denosumab versus
  zoledronic acid for treatment of bone metastases in men with castration-resistant
  prostate cancer: a randomised, double-blind study. *Lancet* 2011;**377**:813-22.
- Henry DH, Costa L, Goldwasser F, Hirsh V, Hungria V, Prausova J et al. Randomized,
  double-blind study of denosumab versus zoledronic Acid in the treatment of bone
  metastases in patients with advanced cancer (excluding breast and prostate cancer) or
  multiple myeloma. *J Clin Oncol* 2011;**29**:1125-32.
- Stopeck AT, Lipton A, Body JJ, Steger GG, Tonkin K, de Boer RH et al. Denosumab
  compared with zoledronic acid for the treatment of bone metastases in patients with
  advanced breast cancer: a randomized, double-blind study. *J Clin Oncol*2010;**28**:5132-9.
- Moher D, Liberati A, Tetzlaff J, Altman DG, PRISMA Group. Preferred reporting items
  for systematic reviews and meta-analyses: the PRISMA statement. *PLoS Med*2009;6:e1000097.
- Royle P, Ford,J, Cummins,E, Mulatero,C. Denosumab for the treatment of bone metastases from solid tumours and multiple myeloma [document on the Internet].
  London: National Institute for Health and Clinical Excellence; 2011 [accessed April 2012]. Available from URL:
  http://guidance.nice.org.uk/TA/Wave21/6/FinalProtocol/pdf/English.
- Higgins JPT, Altman,DG, Sterne,JAC, Cochrane Statistical Methods Group, Cochrane
  Bias Methods Group. *Chapter 8: Assessing risk of bias in included studies [document on the Internet]*. The Cochrane Collaboration; 2011 [accessed August 2011]. Available
  from URL: <u>http://www.cochrane-handbook.org/</u>.

- Lu G, Ades AE. Combination of direct and indirect evidence in mixed treatment
   comparisons. *Stat Med* 2004;**23**:3105-24.
- Andersen PK, Gill RD. Cox's Regression Model for Counting Processes: A Large
   Sample Study. *Ann Stat* 1982;**10**:1100-20.
- 14 Dias S, Welton,NJ, Sutton,AJ, Ades,AE. *NICE DSU Technical Support Document 2: A*general linear modelling framework for pair-wise and network meta-analysis of
  randomised controlled trials. Sheffield: NICE Decision Support Unit; 2011 [accessed
  April 2012]. Available from URL: <u>http://www.nicedsu.org.uk/TSD2%20ES%20-</u>
  <u>%2005\_05\_11\_FINAL\_updated%20Aug2011.pdf</u>.
- Tierney JF, Stewart LA, Ghersi D, Burdett S, Sydes MR. Practical methods for
   incorporating summary time-to-event data into meta-analysis. *Trials* 2007;8:16.
- Rosen LS, Gordon D, Kaminski M, Howell A, Belch A, Mackey J et al. Long-term
  efficacy and safety of zoledronic acid compared with pamidronate disodium in the
  treatment of skeletal complications in patients with advanced multiple myeloma or
  breast carcinoma: a randomized, double-blind, multicenter, comparative trial. *Cancer*2003;**98**:1735-44.
- Kohno N, Aogi K, Minami H, Nakamura S, Asaga T, Iino Y et al. Zoledronic acid
  significantly reduces skeletal complications compared with placebo in Japanese
  women with bone metastases from breast cancer: a randomized, placebo-controlled
  trial. *J Clin Oncol* 2005;**23**:3314-21.
- Lipton A, Theriault RL, Hortobagyi GN, Simeone J, Knight RD, Mellars K et al.
  Pamidronate prevents skeletal complications and is effective palliative treatment in
  women with breast carcinoma and osteolytic bone metastases: long term follow-up of
  two randomized, placebo-controlled trials. *Cancer* 2000;**88**:1082-90.
- Saad F, Gleason DM, Murray R, Tchekmedyian S, Venner P, Lacombe L et al. A
  randomized, placebo-controlled trial of zoledronic acid in patients with hormonerefractory metastatic prostate carcinoma. *J Natl Cancer Inst* 2002;**94**:1458-68.
- Rosen LS, Gordon D, Tchekmedyian S, Yanagihara R, Hirsh V, Krzakowski M et al.
  Zoledronic acid versus placebo in the treatment of skeletal metastases in patients with
  lung cancer and other solid tumors: a phase III, double-blind, randomized trial--the
  Zoledronic Acid Lung Cancer and Other Solid Tumors Study Group. *J Clin Oncol*2003;**21**:3150-7.
- von Moos R, Patrick D, Fallowfield L, Cleeland CS, Henry DH, Qian Y et al. Effects of
  denosumab versus zoledronic acid (ZA) on pain in patients (pts) with advanced cancer
  (excluding breast and prostate) or multiple myeloma (MM): Results from a randomized
  phase III clinical trial. *J Clin Oncol* 2010;**28(Suppl)**:abstr 9043.
- 476 22 Brown JE, Cleeland CS, Fallowfield LJ, Patrick DL, Fizazi K, Smith MR et al. Pain
  477 Outcomes in Patients with Bone Metastases from Castrate-Resistant Prostate Cancer:
  478 Results from A Phase 3 Trial of Denosumab Vs. Zoledronic Acid. *Eur Urol Suppl*479 2011;**10**:336.
- Fallowfield L, Stebbing J, Braybrooke J, Langridge C, Jenkins V. The preferences and
   experiences of different bisphosphonate treatments in women with breast cancer.
   *Psycho-Oncology* 2011;**20**:755-61.

- 483 24 Henry DH, von Moos R, Hungria V, Costa L, Woll PJ, Scagliotti G et al. Delaying
  484 skeletal-related events in a randomized phase III study of denosumab versus
  485 zoledronic acid in patients with advanced cancer. *J Clin Oncol* 2010;**15(Suppl)**:abstr
  486 9133.
- 487 25 Glenny AM, Altman DG, Song F, Sakarovitch C, Deeks JJ, D'Amico R et al. Indirect
   488 comparisons of competing interventions. *Health Technol Assess* 2005;**9**:1-134.
- Cook RJ, Lawless JF. Marginal analysis of recurrent events and a terminating event.
   *Stat Med* 1997;**16**:911-24.
- 491 27 Ford JA, Mowatt G, Jones R. Assessing pharmacological interventions for bone
   492 metastases: the need for more patient-centered outcomes. *Expert Review in Clinical* 493 *Pharmacology* 2012;**5**:271-279
- Stopeck A, Rader M, Henry D, Danese M, Halperin M, Cong Z et al. Cost-effectiveness
  of denosumab vs zoledronic acid for prevention of skeletal-related events in patients
  with solid tumors and bone metastases in the United States [in press]. *J Med Econ*2012;doi: 10.3111/13696998.2012.675380.
- Xie J, Namjoshi M, Wu EQ, Parikh K, Diener M, Yu AP et al. Economic evaluation of
  denosumab compared with zoledronic Acid in hormone-refractory prostate cancer
  patients with bone metastases. *J Manag Care Pharm* 2011;**17**:621-43.
- 30 Body JJ, Diel IJ, Lichinitzer M, Lazarev A, Pecherstorfer M, Bell R et al. Oral
   ibandronate reduces the risk of skeletal complications in breast cancer patients with
   metastatic bone disease: results from two randomised, placebo-controlled phase III
   studies. *Br J Cancer* 2004;**90**:1133-7.
- Body JJ, Diel IJ, Bell R, Pecherstorfer M, Lichinitser MR, Lazarev AF et al. Oral
   ibandronate improves bone pain and preserves quality of life in patients with skeletal
   metastases due to breast cancer. *Pain* 2004;**111**:306-12.
- Tripathy D, Lichinitzer M, Lazarev A, MacLachlan SA, Apffelstaedt J, Budde M et al.
  Oral ibandronate for the treatment of metastatic bone disease in breast cancer:
  efficacy and safety results from a randomized, double-blind, placebo-controlled trial. *Ann Oncol* 2004;**15**:743-50.
- S12 33 Body J-J. Effectiveness and cost of bisphosphonate therapy in tumor bone disease.
   S13 *Cancer* 2003;**97**:859-65.
- 514 34 Diel IJ, Body JJ, Lichinitser MR, Kreuser ED, Dornoff W, Gorbunova VA et al. Improved 515 quality of life after long-term treatment with the bisphosphonate ibandronate in patients 516 with metastatic bone disease due to breast cancer. *Eur J Cancer* 2004;**40**:1704-12.
- Heras P, Kritikos K, Hatzopoulos A, Georgopoulou AP. Efficacy of ibandronate for the
   treatment of skeletal events in patients with metastatic breast cancer. *Eur J Cancer Care (Engl)* 2009;**18**:653-6.
- 520 36 Elomaa I, Blomqvist C, Porkka L, Holmström T, Taube T, Lamberg-Allardt C et al.
   521 Clodronate for osteolytic metastases due to breast cancer. *Biomed Pharmacother* 522 1988;**42**:111-6.

- 37 Paterson AH, Powles TJ, Kanis JA, McCloskey E, Hanson J, Ashley S. Double-blind
   controlled trial of oral clodronate in patients with bone metastases from breast cancer.
   *J Clin Oncol* 1993;**11**:59-65.
- Kristensen B, Ejlertsen B, Groenvold M, Hein S, Loft H, Mouridsen HT. Oral clodronate
   in breast cancer patients with bone metastases: a randomized study. *J Intern Med* 1999;**246**:67-74.
- 39 Dearnaley DP, Sydes MR, Mason MD, Stott M, Powell CS, Robinson AC et al. A
  double-blind, placebo-controlled, randomized trial of oral sodium clodronate for
  metastatic prostate cancer (MRC PR05 Trial). *J Natl Cancer Inst* 2003;**95**:1300-11.
- 40 Elomaa I, Kylmala T, Tammela T, Viitanen J, Ottelin J, Ruutu M et al. Effect of oral
   clodronate on bone pain. A controlled study in patients with metastic prostatic cancer.
   *Int Urol Nephrol* 1992;**24**:159-66.
- Kylmala T, Tammela T, Risteli L, Risteli J, Taube T, Elomaa I. Evaluation of the effect
  of oral clodronate on skeletal metastases with type 1 collagen metabolites. A controlled
  trial of the Finnish Prostate Cancer Group. *Eur J Cancer* 1993;**29A**:821-5.
- 42 Ernst DS, Tannock IF, Winquist EW, Venner PM, Reyno L, Moore MJ et al.
  Randomized, double-blind, controlled trial of mitoxantrone/prednisone and clodronate
  versus mitoxantrone/prednisone and placebo in patients with hormone-refractory
  prostate cancer and pain. *J Clin Oncol* 2003;**21**:3335-42.
- Adami S, Mian M. Clodronate therapy of metastatic bone disease in patients with
   prostatic carcinoma. *Recent Results Cancer Res* 1989;**116**:67-72.
- Adami S, Salvagno G, Guarrera G. Dichloromethylene-diphosphonate in patients with prostatic carcinoma metastatic to the skeleton. *J Urol* 1985;**134**:1152-4.
- 546 45 Kylmala T, Taube T, Tammela TL, Risteli L, Risteli J, Elomaa I. Concomitant i.v. and
  547 oral clodronate in the relief of bone pain--a double-blind placebo-controlled study in
  548 patients with prostate cancer. *Br J Cancer* 1997;**76**:939-42.
- 549 46 Strang P, Nilsson S, Brandstedt S, Sehlin J, Borghede G, Varenhorst E et al. The
  550 analgesic efficacy of clodronate compared with placebo in patients with painful bone
  551 metastases from prostatic cancer. *Anticancer Res* 1997;**17**:4717-21.
- Small EJ, Smith MR, Seaman JJ, Petrone S, Kowalski MO. Combined analysis of two
  multicenter, randomized, placebo-controlled studies of pamidronate disodium for the
  palliation of bone pain in men with metastatic prostate cancer. *J Clin Oncol*2003;**21**:4277-84.
- Smith JA, Jr. Palliation of painful bone metastases from prostate cancer using sodium
  etidronate: results of a randomized, prospective, double-blind, placebo-controlled
  study. *J Urol* 1989;**141**:85-7.
- Arican A, Icli F, Akbulut H, Cakir M, Sencan O, Samur M et al. The effect of two
  different doses of oral clodronate on pain in patients with bone metastases. *Med Oncol*1999;**16**:204-10.
- 50 Brown JE, McCloskey EV, Dewar JA, Body JJ, Cameron DA, Harnett AN et al. The use 563 of bone markers in a 6-week study to assess the efficacy of oral clodronate in patients 564 with metastatic bone disease. *Calcif Tissue Int* 2007;**81**:341-51.

- 565 51 O'Rourke N, McCloskey E, Houghton F, Huss H, Kanis JA. Double-blind, placebo 566 controlled, dose-response trial of oral clodronate in patients with bone metastases. J
   567 Clin Oncol 1995;13:929-34.
- 568 52 Piga A, Bracci R, Ferretti B, Sandri P, Nortilli R, Acito L et al. A double blind
   569 randomized study of oral clodronate in the treatment of bone metastases from tumors
   570 poorly responsive to chemotherapy. *J Exp Clin Cancer Res* 1998;**17**:213-7.
- 571 53 Robertson AG, Reed NS, Ralston SH. Effect of oral clodronate on metastatic bone 572 pain: a double-blind, placebo-controlled study. *J Clin Oncol* 1995;**13**:2427-30.
- 54 Jagdev SP, Purohit P, Heatley S, Herling C, Coleman RE. Comparison of the effects of
  intravenous pamidronate and oral clodronate on symptoms and bone resorption in
  patients with metastatic bone disease. *Ann Oncol* 2001;**12**:1433-8.
- 576 55 Mystakidou K, Stathopoulou E, Parpa E, Kouloulias V, Kouskouni E, Vlahos L. Oral
   577 versus intravenous ibandronic acid: a comparison of treatment options for metastatic
   578 bone disease. *J Cancer Res Clin Oncol* 2008;**134**:1303-10.
- 56 Heras R, I, Zubillaga R, I, Castrillo TM, Montalvo Moreno JJ. Osteonecrosis of the jaws
   and bisphosphonates. Report of fifteen cases. Therapeutic recommendations. *Med* 581 *Oral Patol Oral Cir Bucal* 2007;12:E267-E271.
- 582 57 Berenson JR, Rosen LS, Howell A, Porter L, Coleman RE, Morley W et al. Zoledronic
   583 acid reduces skeletal-related events in patients with osteolytic metastases. *Cancer* 584 2001;**91**:1191-200.
- 585 58 Zaghloul MS, Boutrus R, El-Hossieny H, Kader YA, El-Attar I, Nazmy M. A prospective,
   586 randomized, placebo-controlled trial of zoledronic acid in bony metastatic bladder
   587 cancer. *Int J Clin Oncol* 2010;**15**:382-9.
- 588 59 Zhao YY, Xue C, Hou X, Liao H, Li S, Zhao HY et al. Changes of bone resorption
   589 marker (NTX) in chemotherapy plus zoledronic acid versus chemotherapy alone for
   590 nasopharyngeal cancer patients with bone metastases. *Eur J Cancer* 2011;**47**:848-53.
- Buchali K, Correns HJ, Schuerer M, Schnorr D, Lips H, Sydow K. Results of a double
   blind study of 89-strontium therapy of skeletal metastases of prostatic carcinoma. *Eur J Nucl Med* 1988;**14**:349-51.
- 594 61 Nilsson S, Strang P, Ginman C, Zimmermann R, Edgren M, Nordstrom B et al.
  595 Palliation of bone pain in prostate cancer using chemotherapy and strontium-89. A
  596 randomized phase II study. *J Pain Symptom Manage* 2005;**29**:352-7.
- Porter AT, McEwan AJ, Powe JE, Reid R, McGowan DG, Lukka H et al. Results of a
  randomized phase-III trial to evaluate the efficacy of strontium-89 adjuvant to local field
  external beam irradiation in the management of endocrine resistant metastatic prostate
  cancer. *Int J Radiat Oncol Biol Phys* 1993;**25**:805-13.
- 63 Quilty PM, Kirk D, Bolger JJ, Dearnaley DP, Lewington VJ, Mason MD et al. A
   comparison of the palliative effects of strontium-89 and external beam radiotherapy in
   metastatic prostate cancer. *Radiother Oncol* 1994;**31**:33-40.
- 604 64 Hortobagyi GN, Theriault RL, Porter L, Blayney D, Lipton A, Sinoff C et al. Efficacy of 605 pamidronate in reducing skeletal complications in patients with breast cancer and lytic

- 606 bone metastases. Protocol 19 Aredia Breast Cancer Study Group. *N Engl J Med* 607 1996;**335**:1785-91.
- 608 65 Hortobagyi GN, Theriault RL, Lipton A, Porter L, Blayney D, Sinoff C et al. Long-term
  609 prevention of skeletal complications of metastatic breast cancer with pamidronate.
  610 Protocol 19 Aredia Breast Cancer Study Group. *J Clin Oncol* 1998;**16**:2038-44.
- 611 66 Theriault RL, Lipton A, Hortobagyi GN, Leff R, Gluck S, Stewart JF et al. Pamidronate
  612 reduces skeletal morbidity in women with advanced breast cancer and lytic bone
  613 lesions: a randomized, placebo-controlled trial. Protocol 18 Aredia Breast Cancer
  614 Study Group. J Clin Oncol 1999;17:846-54.
- 67 Rosen LS, Gordon D, Kaminski M, Howell A, Belch A, Mackey J et al. Zoledronic acid
  616 versus pamidronate in the treatment of skeletal metastases in patients with breast
  617 cancer or osteolytic lesions of multiple myeloma: a phase III, double-blind, comparative
  618 trial. *Cancer J* 2001;**7**:377-87.
- 68 Rosen LS, Gordon DH, Dugan W, Jr., Major P, Eisenberg PD, Provencher L et al.
  620 Zoledronic acid is superior to pamidronate for the treatment of bone metastases in
  621 breast carcinoma patients with at least one osteolytic lesion. *Cancer* 2004;**100**:36-43.
- 69 Fallowfield L, Patrick D, Body JJ, Lipton A, Tonkin KS, Qian Y et al. The effect of
  623 treatment with denosumab or zoledronic acid on health-related quality of life in patients
  624 with metastatic breast cancer. *Cancer Research* 2010;**70(Suppl. 2)**:abstr P1-13-05.
- Fallowfield L, Patrick D, Body J, Lipton A, Tonkin KS, Qian Y et al. Effects of
  denosumab versus zoledronic acid (ZA) on health-related quality of life (HRQL) in
  metastatic breast cancer: Results from a randomized phase III trial. *J Clin Oncol*2010;**28(Suppl)**:abstr 1025.
- Martin M, Steger G, von Moos R, Stopeck A, de Boer R, Bourgeois H et al. Benefit of
   denosumab therapy in patients with bone metastases from breast cancer: A number needed-to-treat (NNT) analysis. *Breast* 2011;**20**:S85.
- Stopeck A, Lipton AA, Campbell-Baird C, von Moos R, Fan M, Haddock B et al. Acutephase reactions following treatment with zoledronic acid or denosumab: Results from a
  randomized, controlled phase 3 study in patients with breast cancer and bone
  metastases. *Cancer Research* 2010;**70(Suppl. 2)**:Aastr P6-14-09.
- Stopeck A, Martin M, Ritchie D, Body JJ, Paterson A, Viniegra M et al. Effect of
  denosumab versus zoledronic acid treatment in patients with breast cancer and bone
  metastases: Results from the extended blinded treatment phase. *Cancer Research*2010;**70(Suppl. 2)**:abstr P6-14-01.
- 540 74 Stopeck A, Fallowfield,L, Patrick,D, Cleeland,CS, de Boer,RH, Steger,GG et al. Pain in 541 patients (pts) with metastatic breast cancer: Results from a phase III trial of 542 denosumab versus zoledronic acid (ZA) [document on the Internet]. 33rd Annual San 543 Antiono Breast Cancer Symposium, 8-12 December; 2010 [accessed September 544 2011]. Available from URL: 545 <u>http://www.asco.org/ASCOv2/Meetings/Abstracts?&vmview=abst\_detail\_view&confID=</u> 546 100&abstractID=60225.
- 547 75 Stopeck A, Fallowfield L, Patrick D, Cleeland CS, de Boer RH, Steger GG et al. Effects
   of denosumab versus zoledronic acid (ZA) on pain in patients (pts) with metastatic

- breast cancer: Results from a phase III clinical trial. *J Clin Oncol* 2010;**28(Suppl)**:abstr
  1024.
- 651 76 Saad F, Olsson C, Schulman CC. Skeletal morbidity in men with prostate cancer:
  652 quality-of-life considerations throughout the continuum of care. *Eur Urol* 2004;**46**:731653 9.
- Saad F, Gleason DM, Murray R, Tchekmedyian S, Venner P, Lacombe L et al. Longterm efficacy of zoledronic acid for the prevention of skeletal complications in patients
  with metastatic hormone-refractory prostate cancer. *J Natl Cancer Inst* 2004;**96**:87982.
- Saad F. Clinical benefit of zoledronic acid for the prevention of skeletal complications
   in advanced prostate cancer. *Clin Prostate Cancer* 2005;**4**:31-7.
- Saad F, Lipton A, Cook R, Chen YM, Smith M, Coleman R. Pathologic fractures
   correlate with reduced survival in patients with malignant bone disease. *Cancer* 2007;**110**:1860-7.
- 80 Saad F, Chen YM, Gleason DM, Chin J. Continuing benefit of zoledronic acid in
   preventing skeletal complications in patients with bone metastases. *Clin Genitourin Cancer* 2007;**5**:390-6.
- 666 81 Saad F, Eastham J. Zoledronic Acid improves clinical outcomes when administered 667 before onset of bone pain in patients with prostate cancer. *Urol* 2010;**76**:1175-81.
- Weinfurt KP, Anstrom KJ, Castel LD, Schulman KA, Saad F. Effect of zoledronic acid
   on pain associated with bone metastasis in patients with prostate cancer. *Ann Oncol* 2006;**17**:986-9.
- 83 Rosen LS, Gordon D, Tchekmedyian NS, Yanagihara R, Hirsh V, Krzakowski M et al.
  Long-term efficacy and safety of zoledronic acid in the treatment of skeletal
  metastases in patients with nonsmall cell lung carcinoma and other solid tumors: a
  randomized, Phase III, double-blind, placebo-controlled trial. *Cancer* 2004;**100**:261321.
- 84 Schulman CC. Efficacy of zoledronic acid in the treatment of bone metastases
  secondary to renal cell carcinoma. *Eur Urol Suppl* 2004;**3**:40-5.

679 Table 1: Studies meeting inclusion criteria but unsuitable for NMA

Primary tumour	Study ID	Intervention	Comparator	Reason for exclusion
	cebo/ another BP			
(n=27)				
Breast	Body 2004 <sup>30</sup> Body 2004 <sup>31</sup> Tripathy 2004 <sup>32</sup>	Ibandronate (oral)	Placebo	SRE definition not comparable
	Body 2003, <sup>33</sup> Diel 2004 <sup>34</sup>	Ibandronate (iv)	Placebo	SRE definition not comparable
	Heras 2009 <sup>35</sup>	Ibandronate (iv)	Placebo	SRE definition not comparable
	Elomaa 1988 <sup>36</sup>	Clodronate (oral)	Placebo	SRE definition not comparable
	Paterson 1993 <sup>37</sup>	Clodronate (oral)	Placebo	SRE definition not comparable
	Kristensen 1999 <sup>38</sup>	Clodronate (oral)	Open	SRE definition not comparable
Prostate	Dearnaley 2003 <sup>39</sup>	Clodronate (oral)	Placebo	Hormone sensitive prostate cancer
	Elomaa 1992 <sup>40</sup>	Clodronate (iv)	Placebo	Only painful metastases
	Kylmala 199341	Clodronate (iv)	Open	Only painful metastases
	Ernst 2003 <sup>42</sup>	Clodronate (iv)	Placebo	Unlicensed administration of clodronate
	Adami 1989, <sup>43</sup> Adami 1985 <sup>44</sup>	Clodronate (iv+im+oral)	Placebo	Only painful metastases
	Kylmala 199745	Clodronate (iv+oral)	Placebo	Only painful metastases
	Strang 1997 <sup>46</sup>	Clodronate (iv)	Placebo	Only painful metastases
	Small 2003 <sup>47</sup>	Pamidronate (iv)	Placebo	Only painful metastases
	Smith 1989 <sup>48</sup>	Etidronate (iv+oral)	Placebo	Only painful metastases
OST	Arican 1999 <sup>49</sup>	Clodronate (oral)	Placebo	SRE definition not comparable
	Brown 2007 <sup>50</sup>	Clodronate (oral)	Placebo	Outcomes not relevant
	O'Rourke 1995 <sup>51</sup>	Clodronate (oral)	Placebo	Outcomes not relevant
	Piga 1998 <sup>52</sup>	Clodronate (oral)	Placebo	Outcomes not relevant
	Robertson 1995 <sup>53</sup>	Clodronate (oral)	Placebo	SRE definition not comparable
	Jagdev 2001 <sup>54</sup>	Clodronate (oral)	Pamidronate (iv)	Outcomes not relevant
	Mystakidou 2008 <sup>55</sup>	Ibandronate (oral)	Ibandronate (iv)	Outcomes not relevant
	Heras 2007 <sup>56</sup>	Ibandronate (iv)	Placebo	SRE definition not comparable
	Berenson 2001 <sup>57</sup>	Zoledronic acid (iv)	Pamidronate (iv)	SRE definition not comparable
	Zaghloul 2010 <sup>58</sup>	Zoledronic acid (iv)	Placebo	SRE definition not comparable
	Zhao 2011 <sup>59</sup>	Zoledronic acid (iv)	Open	SRE definition not comparable

Primary tumour	Study ID	Intervention	Comparator	Reason for exclusion				
BSC vs placebo/ another BSC								
(n=4)								
Prostate	Buchali 1988 <sup>60</sup>	Strontium chloride (iv)	Placebo	SRE definition not comparable				
	Nilsson 2005 <sup>61</sup>	Strontium chloride (iv)	FEM	Only painful metastases				
	Porter 1993 <sup>62</sup>	Strontium chloride (iv)	Placebo	Only painful metastases				
	Quilty 1994 <sup>63</sup>	Strontium chloride (iv)	Radiotherapy	Only painful metastases				

#### 682 Table 2: Characteristics of studies included in NMA

Author, year, countryCancer typeInter sand durationInter		Intervention	Participants		Outcomes	Comments
		5	Age	Prev SRE, n (%)		
Kohno 2005 <sup>17</sup> Country: Japan	Breast	Zoledronic acid 4 mg (n=114)	mean 54.3	39 (34.2)	SRE outcomes <i>Ratio of SRE rate</i> (defined as the total number of SREs divided by the total years on study) for patients treated with zoledronic acid divided by the SRE rate for the placebo group (excluding HCM in	Both administered via 15-minute infusion.
Duration : 12 months		Placebo (n=113)	mean 53.5	47 (41.6)	definition) Proportion of patients experiencing at least one SRE Time to first SRE Multiple-event analysis by the Andersen-Gill method Risk ratio for developing SREs Other outcomes Change from baseline BPI composite pain scores and bone resorption markers	Infusions were administered every 4 weeks for 12 months
Lipton 2000 <sup>18,64-</sup> 66 Country:	Breast	Pamidronate 90 mg (n=367)	<50 years 25% 51-65 years 42% >65 years 33%	NR	SRE outcomes <i>SMR</i> (number of skeletal complications per time on trial for each patient (events/year); the overall SMR was calculated with and without hypercalcemia counted as a skeletal complication	Both administered in 250 mL of 5% dextrose in
US Duration : 24 months (24 cycles)		Placebo (n=384)	<pre>&lt;50 years 33% &lt;50 years 29% 51-65 years 38% &gt;65 years 34%</pre>	NR	Proportion of patient with skeletal complications Time from randomisation to first SRE Other outcomes Bone pain score, analgesic use, ECOG performance status and quality of life measured as mean change from baseline to 24 months or last visit (any time during study); Overall survival	water given as a 2-hour intravenous infusion every 3- 4 weeks for 24 cycles.
Rosen 2003a <sup>16,6</sup> <sup>7,68</sup> Country:	Breast cancer	Zoledronic acid 4 mg (n=378)	median 58	232 (61.4)	SRE outcomes Proportion of patients who experienced at least 1 SRE during 25 month study period (HCM not included). Proportion of patients experiencing any SRE (including HCM)	Both administered as an intravenous infusion
Multinati onal Duration : 25		Pamidronate 90 mg (n=388)	median 56	244 (62.9)	Time to first SRE SMR* Multiple-event analysis* Other outcomes	depending on the scheduling of other antineoplastic

months					None reported	treatments every 3–4 weeks for 24 months
Stopeck 2010 <sup>8,69-</sup> 75 Country: Multinati onal Duration : 34 months	Breast	Denosumab 120 mg (subcutaneo us injection) + placebo (intravenous infusion) (n=1026) Zoledronic acid 4 mg (intravenous infusion) + placebo (subcutaneo us injection) (n=1020)	mean 57 mean 56	378 (36.8)	SRE outcomes Time to first on-study SRE (non-inferiority test) Time to first on-study SRE (superiority test) Time to first and subsequent on-study SREs (multiple event analysis). [Subsequent events must have occurred at least 21 days apart from the most recent event to ensure that linked events (eg, surgery to repair a fracture or multiple doses of radiation during a course of treatment) were not counted as separate SREs.] Other outcomes Overall survival Disease progression Skeletal morbidity rate Percent change in uNTx and BSAP levels.	Intravenous products (placebo or zoledronic acid) were dose- adjusted on the basis of baseline creatinine clearance 60 mL/min and were held for renal function deterioration on- study as per zoledronic acid prescribing
Fizazi 2011 <sup>6</sup> Country: Multinati onal Duration : 27 months	Prostate	Denosumab 120 mg (subcutaneo us) + placebo (n=950) Zoledronic acid 4 mg + placebo (subcutaneo us) (n=951)	median 71 median 71	232 (24)	SRE outcomes Time to first on-study skeletal-related event; assessed for non- inferiority If testing of the primary endpoint showed non-inferiority, then the same outcome was further tested as a secondary endpoint, together with the secondary endpoint of time to first and subsequent on-study skeletal-related events (multiple events), for superiority Other outcomes Overall survival Overall disease progression Prostate-specific antigen concentration during the study Change in bone turnover markers from baseline Pain	information Interventions given every 4 weeks until the primary analysis cut off date. Dose adjustment as per Stopeck 2010
Saad 2002 <sup>19,76-</sup> 82 Country: Multinati onal	Prostate	Zolendronic acid 4mg (n=214) Placebo	mean 72 mean 72	66 (30.8) 78 (37.5)	SRE outcomes The proportion of patients having at least one skeletal-related event Time to the first skeletal- related event Skeletal morbidity rate Proportion of patients with individual skeletal-related events Other outcomes	Administered every 3 weeks for 15 months (20 cycles). Initially 5 min infusion (in

Duration : 15 months		(n=208)		110 (50)	Time to disease progression Objective bone lesion response Bone biochemical markers Quality-of-life parameters Pain	50ml), changed to 15 min infusion (in 100ml) in 1999
Henry 2011 <sup>7,21,2</sup> 4 Country:	Other solid tumours	<b>Denosumab</b> <b>120 mg</b> (n=890)	median 61	446 (50)	SRE outcomes <i>Time to first on-study SRE (non-inferiority)</i> Time to first on-study SRE (superiority tests) Time to first-and-subsequent SRE (multiple-event analysis).	Zoledronic acid administered intravenously monthly with
Multinati onal Duration : 7 months (median time on- study		Zoledronic acid 4 mg (n=886)	median 60	440 (50)	Other outcomes Bone turnover markers Overall survival Overall disease progression.	subcutaneous placebo.
Rosen 2003b <sup>20,8</sup> <sup>3,84</sup> Country:	Other solid tumours	Zoledronic acid 4 mg (n=257)	median 64	166 (65)	SRE outcomes Proportion of patients with at least one SRE Time to first SRE SMR (defined as the number of SREs per year) Multiple event	Interventions administered intravenously every 3 weeks
Multinati onal Duration of study: 9 months		Placebo (n=250)	median 64	179 (73)	analysis <b>Other outcomes</b> Pain score Analgesic use ECOG performance status Best bone lesion response and time to progression of bone lesions Biochemical markers of bone resorption Time to progression of overall disease and survival. Quality of life	for 9 months

Cancer	Study	Intervention	TTF SRE		p value	TTF+S SRE	SMR	p value
Breast	Kohno 2005 <sup>17</sup>	Zoledronic acid (n=114)	Not reached	N/R	0.007	RR 0.59 (0.38 to 0.91)	0.63	0.016
		Placebo (n=113)	364 days (~12.1 months)	1			1.1	
	Lipton 2000 <sup>18</sup>	Pamidronate (n=367)	12.7 months (95%CI 9.6 to 17.2)	N/R	<0.001	NR	2.4 (5.5)	<0.001
		Placebo (n=387)	7.0 months (95%CI 6.2 to 8.5)				3.7 (5.5)	
	Rosen	Zoledronic acid	349 days(chemo treated)	N/R	0.826	RR = 0.80 (0.66	0.9	0.125
2003a <sup>16</sup>	(n=378)	415 days(hormone treated)		(chemo)	to 0.97)		_	
		Pamidronate (n=388)	366 days (chemo treated) 370 days(hormone treated)		0.047 (hormone)		1.49	
	Stopeck 2010 <sup>8</sup>	Denosumab (n=1026)	Not reached	HR 0.82 95%CI 0.71	<0.001	RR *0.77 (0.66 to 0.89)	0.45	0.004
		Zoledronic acid (n=1020)	26.4 months	to 0.95			0.58	
Prostate	Fizazi	Denosumab (n=950)	20.7 months	HR 0.82,	0.0002	RR* 0.82 (95%	NR	NR
	2011 <sup>6</sup>	Zoledronic acid (n=951)	17.1 months	95%CI 0.71 to 0.95		CI 0.71 to 0.94)	NR	
	Saad 2002 <sup>19</sup>	Zoledronic acid (n=214)	361 days (prev SRE) 499 days (no prev SRE)	N/R	0.066 (prev SRE)	RR 0.64 (95% CI not reported, p value 0.002)	0.80	0.006
		Placebo (n=208)	258 days (prev SRE) 337 days (no prev SRE)		0.065 (no prev SRE)	, ,	1.49	
Other	Henry	Denosumab (n=886)	20.6 months	HR 0.84,	0.0007	RR*† 0.90 (0.77	NR	NR
solid tumours	2011 <sup>7</sup>	Zoledronic acid (n=890)	16.3 months	95%CI 0.71 to 0.98		to 1.04)	NR	
	Rosen	Zoledronic acid	230 days	N/R	0.023	HR 0.732,	2.24	0.069
	2003 <sup>20</sup>	Placebo	163 days	]		p=0.017	2.52	

685 Table 3: Results of individual studies included in the NMA

 $RR = risk ratio, RR^* = rate ratio, HR = hazard ratio, † = includes multiple myeloma, N/R = not reported, TTF SRE = time to first skeletal related$ event, TTF+S SRE = time to first and subsequent skeletal related events

# 688 Table 4: Risk of bias of studies included in NMA

Study id	Q1 Adequate sequence generation?	Q2 Adequate allocation concealment?	Q3 Blinding?	Q4 Incomplete outcome data addressed?	Q5 Free of selective reporting?
Breast cancer					
Lipton 2000 <sup>18</sup>	Low	Low	Low	Unclear	Unclear
Kohno 2005 <sup>17</sup>	Low	Low	Low	High	Low
Stopeck 2010 <sup>8</sup>	Unclear	Unclear	Low	Low	Low
Rosen 2003a <sup>16</sup>	Low	Low	Low	Low	Low
Prostate cancer				·	
Fizazi 2011 <sup>6</sup>	Low	Low	Low	Low	Low
Saad 2002 <sup>19</sup>	Low	Low	Low	Low	Low
Other solid tumours				·	
Henry 2011 <sup>7</sup>	Low	Low	Low	Low	Low
Rosen 2003b <sup>20</sup>	Unclear	Unclear	Low	High	Low

Table 5: Breast cancer NMA results

Comparison	TTF SRE	TTF+S Risk	SMR
	HR (95% CI)	Ratio (95% CI)	Rate Ratio (95% CI)
Denosumab versus	0.82 (0.71 to	0.77 (0.66 to	0.90 (0.67 to 1.09)
zoledronic acid	0.95)	0.89)	
Denosumab versus pamidronate	0.79 (0.61 to 1.03)	0.62 (0.48 to 0.80)	0.73 (0.41 to 1.06)
Denosumab versus	0.46 (0.29 to	0.45 (0.28 to	0.47 (0.25 to 0.67)
placebo	0.72)	0.72)	
Zoledronic acid versus	0.56 (0.36 to	0.59 (0.37 to	0.52 (0.32 to 0.70)
placebo	0.86)	0.91)	

TTF SRE = time to first skeletal related event, TTF+S SRE = time to first and subsequent skeletal related events, SMR = skeletal morbidity rate

Table 6: Prostate cancer NMA results

	TTF SRE HR (95%CI)	TTF+S Risk Ratio (95% CI)	SMR Rate Ratio (95% CI)
Denosumab versus	0.82 (0.71 to	0.82 (0.71 to	0.95 (0.46 to 1.47)
zoledronic acid	0.95)	0.94)	
Denosumab versus	0.56 (0.40 to	0.53 (0.39 to	0.52 (0.07 to 0.82)
placebo	0.77)	0.72)	
Zoledronic acid versus	0.68 (0.50 to	0.64 (0.48 to	0.54 (0.11 to 0.83)
placebo	0.91)	0.85)	

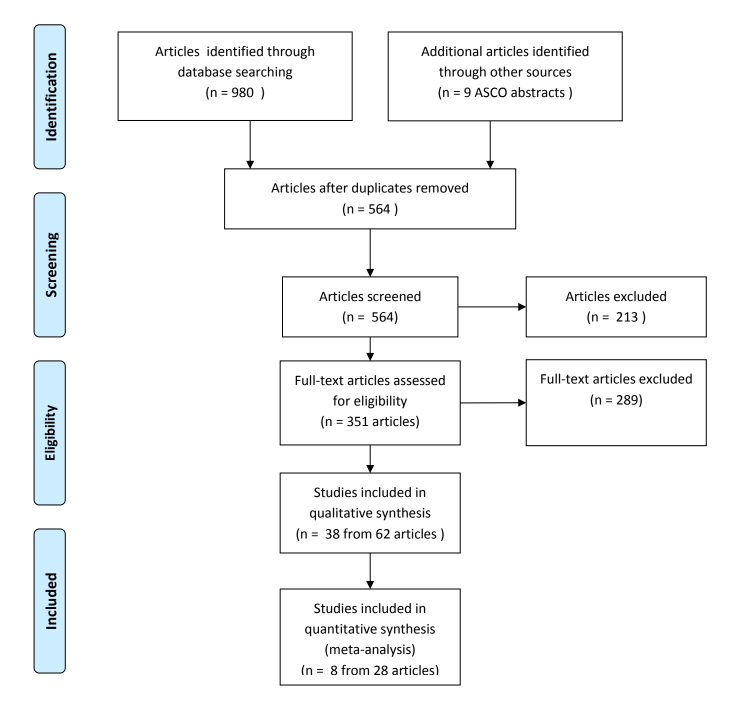
TTF SRE = time to first skeletal related event, TTF+S SRE = time to first and subsequent skeletal related events, SMR = skeletal morbidity rate

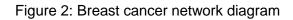
Table 7: Other solid tumours and non small cell lung cancer NMA results

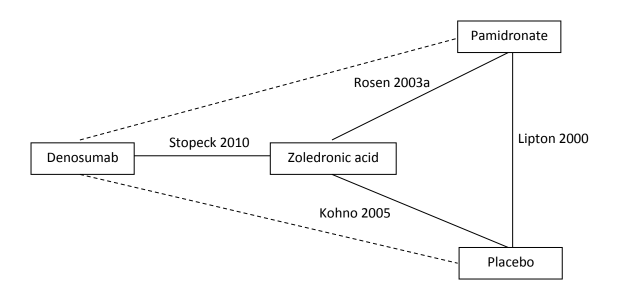
	Other s	olid tumours	1	NSCLC
	TTF SRE	TTF+S SRE	TTF SRE	TTF+S SRE
	HR (95%CI)	RR (95%CI)	HR (95%CI)	RR (95%CI)
Denosumab versus zoledronic	0.79 (0.62 to 0.99)	0.83 (0.67 to 1.03)	0.84 (0.64 to 1.10)	0.87 (0.68 to 1.12)
acid				
Denosumab versus placebo	0.30 (0.11 to 0.82)	0.61 (0.39 to 0.97)	0.68 (0.45 to 1.03)	0.63 (0.42 to 0.97)
Zoledronic acid versus placebo	0.37 (0.14 to 1.01)	0.74 (0.49 to 1.10)	0.81 (0.59 to 1.11)	0.73 (0.52 to 1.02)

TTF SRE = time to first skeletal related event, TTF+S SRE = time to first and subsequent skeletal related events, SMR = skeletal morbidity rate

# Figure 1: PRISMA flow diagram





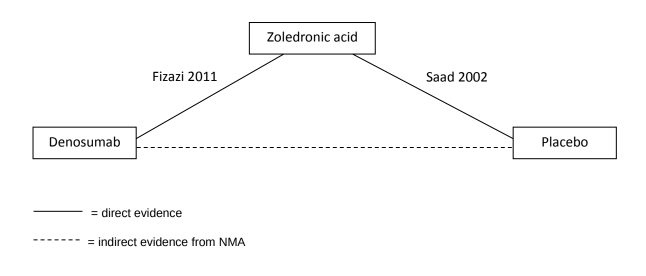


----- = direct evidence

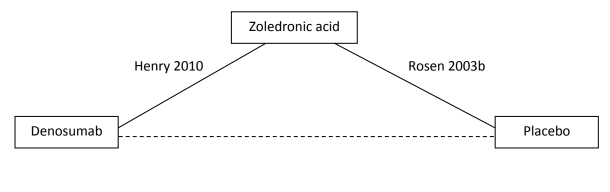
= indirect evidence from NMA

Note: Lipton 2000 data was only available for the SMR outcome.

Figure 3: Prostate cancer network diagram



# Figure 4: Other solid tumours network



----- = direct evidence

== indirect evidence from NMA