Section 1: Introductory statements and governance

COST SAVINGS IMPACT OF BIOSIMILARS: A LOCAL INDIRECT COMPARISON OF BRIGATINIB VERSUS TUZUMAB BIOBASEMILAR

Background and importance

Boosting biosimilars is an indispensable approach in conducting cost savings management in healthcare systems. In fact, biosimilars can provide similar effectiveness and safety to originators but with lower costs, and can increase market competition.

Aim and objectives

The purpose of this paper was to demonstrate the economic advantage of a trastuzumab biosimilar in real practice, showing and comparing costs and consumption during the period 2018–2019.

Material and methods

To conduct this analysis, patients, type of prescription (originator or biosimilar), number of cycles, administered milligrams and purchase prices, during the period 2018–2019, were extrapolated from pharmacy software and matched. A simulation was also performed to estimate potential savings, based on three scenarios of different biosimilar penetration rates (50%, 75% and 100%) and using actual costs.

Results

Compared with 2018, during 2019, the number of treated patients remained similar (102 vs 98) and both a reduction of administered units of trastuzumab originator (TO) (11383) and a growing number of prescriptions of trastuzumab biosimilar (TB) (+833 units) were observed. Costs of TO decreased from 3.28€/mg to 2.58€/mg, while average TB cost was 1.07€/mg in 2019. TB accounted for 24% of trastuzumab prescriptions but with only 11% of total costs, resulting in a reduction in total expenditure of 1 045 540€ spent in 2018 to 1 248 826€ in 2019. In addition, a simulation was performed considering three hypothetical scenarios with different penetration rates of TB in the market share (50%, 75% and 100%) and using actual costs.

Conclusion and relevance

Currently, cost savings and rationalisation policy are playing an essential role in healthcare systems, and biosimilars represent a great opportunity to reallocate available resources. This study demonstrated that enhancing a trastuzumab biosimilar is a good strategy for the sustainability of care. Trastuzumab costs decreased while the number of patients remained similar. This positive result was due to both the introduction of new biosimilars and the reduction of the costs of the originators. In summary, biosimilars constitute an efficient strategy for the sustainability of national health services, allowing resource reallocation and access to care to a larger number of patients.

REFERENCES AND/OR ACKNOWLEDGEMENTS

Conflict of interest

No conflict of interest

Abstract 11S-021

INDIRECT COMPARISON OF BRIGATINIB VERSUS ALECTINIB IN ALK POSITIVE NON-SMALL CELL LUNG CANCER

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10.1136/ejhpharm-2021-eahpconf.2

Background and importance

ALK gene mutation occurs in 3–5% of patients with non-small cell lung cancer (NSCLC). Brigatinib and alectinib are potent ALK tyrosine kinase inhibitors, indicated in NSCLC.

Aim and objectives

The aim of this study was to perform an adjusted indirect treatment comparison (ITC) of the efficacy of brigatinib and alectinib in patients with NSCLC using a common comparator, and to establish whether both ALK inhibitors can be declared equivalent therapeutic alternatives (ETA).

Material and methods

A search was carried out to detect clinical trials (CTs) with brigatinib or alectinib with similar populations, endpoints and follow-up periods. If multiple studies were found for the same drug, the results were combined in a meta-analysis using the Metasurv calculator. ITC was done according to Bucher’s method. To establish the positioning, ETA guidelines were applied. Delta value, maximum acceptable difference as a clinical criterion of non-inferiority, was set at 0.64 (and its inverse, 1.57) (the value used in the calculation of the sample size) for progression free survival (PFS).

Conclusion and relevance

ITC showed no statistically significant differences in PFS between brigatinib and alectinib for Asian and non-Asian patients. The extent of the 95% CI
values showed some uncertainty. According to the ETA guidelines, as the percentage outside the delta margin was small, both drugs could be considered as ETA in most patients with ALK positive NSCLC.

REFERENCES AND/OR ACKNOWLEDGEMENTS

Conflict of interest No conflict of interest

11SG-022 ECONOMIC ANALYSIS AFTER THE INCORPORATION OF BEVACIZUMAB BIOSIMILAR IN A THIRD LEVEL HOSPITAL

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10.1136/ejhpharm-2021-eahpconf.1

Background and importance The introduction of biosimilars after the end of the patenting of biologics is an opportunity for significant savings in pharmaceutical spending, contributing to the sustainability of the health system.

Aim and objectives To estimate the economic savings achieved by switching to bevacizumab biosimilar in a third level hospital.

Material and methods A descriptive observational study was conducted during 2019 to estimate the economic impact of the incorporation of bevacizumab biosimilar, one of the objectives of the Annual Management Plan of our Autonomous Community, Castilla y León. The switch was agreed with the oncology service for all patients, except those who did not meet the criteria for the indication of the drug. Variables collected were: number of patients, total annual cost and quarterly cost variation 2019/2020. To estimate the cost (€), the net unit price was used (PVL-discounts (official and laboratory) + 4% VAT), and the Farmatools application was used to obtain the data.

Results The switch began in July 2020, with the degree of penetration of 96.7% in the oncology service because 2 patients (3.3%) did not switch to the biosimilar because their indication was not included in the technical data sheet. A quarterly cut-off (July–September 2019 vs 2020) showed a similar number of patients treated with original bevacizumab (n=50) and bevacizumab biosimilar (n=58) with a total cost during those months of 194 543€ and 119 001€ respectively. Although the number of patients treated with bevacizumab biosimilar in the third quarter of 2020 was higher than in the previous year with original bevacizumab (eight more patients compared with July–September 2019), the cost was reduced by 38.8%.

In 2019, 121 patients were treated with the original bevacizumab, for a total annual cost of 945 710€ and an annual expense/patient of 7816€. In 2020, assuming a cost reduction of 38.8% and the same number of patients treated as in 2019 (121 patients), annual expenditure is estimated at 578 891€ and a saving of 366 819€ per year.

Conclusion and relevance The introduction of biosimilars is an efficient measure to reduce hospital pharmaceutical expenses, maintaining the same effectiveness and safety as the original medicine, contributing to the sustainability of the National Health System.

References and/or acknowledgements

Conflict of interest No conflict of interest

11SG-023 PHARMACOECONOMIC ANALYSIS OF REFERENCE BEVACIZUMAB: OPPORTUNITY FOR IMPROVED EFFICIENCY

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10.1136/ejhpharm-2021-eahpconf.4

Background and importance The recent approval of bevacizumab biosimilar (Beva-Bs) raises the possibility of a more efficient drug therapy. Reference bevacizumab (Beva-Ref) was the cancer drug with the greatest impact in our health area in 2019.

Aim and objectives To evaluate the pharmacoeconomic impact of Beva-Ref in oncological therapy in 2019 and to analyse measures that promote its therapeutic optimisation, such as more efficient dosage regimens (DR) and implementation of Beva-Bs.

Material and methods This was a descriptive retrospective study made in a level II hospital. Farhos-v5.3.3 was used as the pharmacotherapeutic management tool for cancer patients treated with Beva-Ref during 2019. Economic data were collected from the Gestión–Farmatools module.

- Pharmacoeconomic analysis was done by therapeutic cost of Beva-Ref use in 2019. Therefore, cost/indication consumption and therapeutic scheme were recorded.
- Therapeutic optimisation measures analyses were conducted according to efficient DR, in concordance with the product monograph.
- Possibility of using Beva-Bs: hypothetical savings were estimated on 2019’s annual consumption, assuming switching to Beva-Bs: (a) 100% of patients; (b) only new patients.

Results 58 patients were treated in 2019. Total cost was 710 842€ and according to indication: nine breast cancer 210 106€ (30%); 25 metastatic colorectal cancer (mCRC) 205 671€ (29%); and 11 ovarian cancer 165 346€ (23%). 41 patients (71%) started treatment with a total cost of 406 897€, mostly: mCRC 169 274€ (42%); 4 breast cancer 74 139€ (18%); and 7 ovarian cancer 65 776€ (16%). Treatment continuations: 17 patients (29%) at a cost of 303 943€, mainly 5 breast cancer 135 967€ (45%), 4 ovarian cancer 99 570€ (33%) and 4 mCRC 36 396€ (12%). The most efficient DR in mCRC was prescribed 100%. In the remaining diagnoses, DR was achieved, except for ovarian/ endometrial cancer, with agreement of 45% and 0%, respectively.

With respect to the possibility of using Beva-Bs: a saving of 312 800€ was estimated if switching to Beva-BS in all patients, with savings in breast cancer 92 450€, mCRC 90 500€ and ovarian cancer 72 750€. Considering only new patients, savings would be 179 000€, mostly mCRC, breast and ovarian cancer (74 500€, 32 600€ and 28 900€, respectively).

Conclusion and relevance The 2019 results showed efficient DR, and consequently the potential for cost containment, given the incorporation of Beva-Bs into our therapeutic arsenal, and would be key for universal access to the best therapeutic option.

References and/or acknowledgements

Conflict of interest No conflict of interest