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SYSTEMATIC REVIEW



A Systematic Review of the Cost-Effectiveness of Biologics for Ulcerative Colitis

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Abstract

Background Ulcerative colitis (UC) is a chronic autoimmune inflammation of the colon. The condition significantly decreases quality of life and generates a substantial economic burden for healthcare payers, patients and the society in which they live. Some patients require chronic pharmacotherapy, and access to novel biologic drugs might be crucial for long-term remission. The analyses of costeffectiveness for biologic drugs are necessary to assess their efficiency and provide the best available drugs to patients.

Objective Our aim was to collect and assess the quality of economic analyses carried out for biologic agents used in the treatment of UC, as well as to summarize evidence on the drivers of cost-effectiveness and evaluate the transferability and generalizability of conclusions.

Methods A systematic database review was conducted using MEDLINE (via PubMed), EMBASE, Cost-Effectiveness Analysis Registry and CRD0. Both authors independently reviewed the identified articles to determine their eligibility for final review. Hand searching of references in collected papers was also performed to find any relevant articles. The reporting quality of economic analyses included was evaluated by two reviewers using the International Society of Pharmacoeconomics and Outcomes Research (ISPOR) Consolidated Health Economic Evaluation Reporting Standards (CHEERS) statement checklist. We reviewed the sensitivity analyses in costeffectiveness analyses to identify the variables that may have changed the conclusions of the study. Key drivers of cost-effectiveness were selected by identifying uncertain parameters that caused the highest change of the results of the analyses compared with base-case results.

Results Of the 576 identified records, 87 were excluded as duplicates and 16 studies were included in the final review; evaluations for Canada, the UK and Poland were mostly performed. The majority of the evaluations revealed were performed for infliximab (approximately 75% of total volume); however, some assessments were also performed for adalimumab (50%) and golimumab (31%). Only three analyses were conducted for vedolizumab, whereas no relevant studies were found for etrolizumab and tofacitinib. The reporting quality of the included economic analyses was assessed as high, with an average score of 21 points per 24 maximum possible (range 14-23 points according to the ISPOR CHEERS statement checklist). In the case of most analyses, quality-adjusted life-years were used as a clinical outcome, and endpoints such as remission, response and mucosal healing were less common. The higher clinical effectiveness (based on response rates) of biological treatment over non-biological treatments was presented in revealed analyses. The incremental cost-utility ratios for biologics, compared with standard care, varied significantly between the studies and ranged from US\$36,309 to US\$456,979. The lowest value was obtained for infliximab and the highest for the treatment scheme including infliximab 5 mg/kg and infliximab 10 mg/ kg + adalimumab. The change of utility weights and clinical parameters had the most significant influence on the

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results of the analysis; the variable related to surgery was the least sensitive.

Conclusions Limited data on the cost-effectiveness of UC therapy were identified. In the majority of studies, the lack of cost-effectiveness was revealed for biologics, which was associated with their high costs. Clinical outcomes are transferable to other countries and could be generalized; however, cost inputs are country-specific and therefore limit the transferability and generalizability of conclusions. The key drivers and variables that showed the greatest effect on the analysis results were utility weights and clinical parameters.

Key Points

The majority of the identified economic evaluations related to the most commonly used agents: infliximab, adalimumab and golimumab. Most analyses were performed for Canada, the UK and Poland.

Higher clinical effectiveness (reported as response rates) of biologic drugs over reference therapies was revealed and additional clinical effect of biologics in terms of quality-adjusted life-years was presented in economic analyses. The cost-effectiveness of biological treatment, compared with standard care alone, was reported in three analyses, and, in nine studies, biologics were described as inefficient.

Infliximab was cost-effective when compared with cyclosporine or surgical treatment.

Utility weights and clinical parameters are the main key drivers of the cost-effectiveness of biologic treatments in UC.

1 Introduction

Ulcerative colitis (UC) is an inflammatory bowel disease characterized by the chronic inflammation of colonic mucosa. Patients with UC typically present with bloody diarrhoea, the passage of pus or mucus (or both), and abdominal cramping during bowel movements [1]. The clinical course is difficult to predict and changeable due to alternating periods of remissions and exacerbations, which may occur spontaneously or in response to external triggers (e.g. treatment changes) [2, 3].

The annual incidence of UC is from 0 to 19.2 per 100,000 persons in North America, and from 0.6 to 24.3 per 100,000 persons in Europe. The prevalence ranges from

37.5 to 248.6 per 100,000 persons in North America, and from 4.9 to 505 per 100,000 persons in Europe [4].

The main goal of UC pharmacotherapy is to effectively obtain the remission of symptoms, and subsequently maintain symptom-free periods. Patients affected with UC often require expensive, lifelong treatment, and sometimes surgery, both of which generate great direct costs to the healthcare payer. Conventional therapy includes corticosteroids, 5-aminosalicylates (5-ASA; 5-aminosalicylic acid) and thiopurines [3]. Biologics, which target a specific inflammatory mediator, are used in patients with moderateto-severe chronic UC who are unresponsive or intolerant to current therapy. The following biologics are currently being assessed or are already used for UC therapy: tumour necrosis factor inhibitors (infliximab, adalimumab, golimumab), integrin antagonists (vedolizumab, etrolizumab) and Janus kinase (JAK) inhibitors (tofacitinib) [5].

Clinical trials revealed that the biologics are effective in UC therapy, but also generate a heavy burden on healthcare service providers and payers [5, 6]. The increasing prevalence and high costs of UC pharmacotherapy and surgery constitute a considerable challenge for healthcare systems and generate a need for efficient and cost-effective management of this chronic long-term disease [7]. The cost-effectiveness of different treatment strategies is required to make objective and verifiable reimbursement decisions, providing patients with the best treatment regimens available.

Previous reviews on a similar topic did not provide a comprehensive assessment of the quality of cost-effectiveness analyses and the key drivers of cost-effectiveness were not identified. The objective of our study was to systematically review medical databases to collect relevant publications on the cost-effectiveness of biologic treatment in UC, and to perform a sophisticated and extensive review of the methodologies used in the economic evaluations, as well as assess the quality of the collected studies and identify the key drivers of cost-effectiveness.

2 Materials and Methods

2.1 Systematic Literature Search

A systematic literature review was performed in July 2017 in order to obtain data on the efficiency of the biologic treatments of UC. The following databases were searched: MEDLINE (via PubMed), EMBASE, the Cost-Effectiveness Analysis (CEA) Registry and Centre for Reviews and Dissemination (CRD). The following terms were used to define the population studied: ulcerative colitis, ulcerative disease, colitis ulcerosa. We focused on the biologics that are authorized for use in UC in Europe and/or in the US: infliximab, adalimumab, golimumab, vedolizumab, etrolizumab, and tofacitinib. The following terms were used to define the intervention: biologic, TNF, tumor necrosis factor, tumour necrosis factor, infliximab, adalimumab, golimumab, Remsima, Remicade, Inflectra, Humira, Simponi, integrin antagonist, vedolizumab, Entyvio, JAK inhibitor, Janus kinases, etrolizumab, tofacitinib, Xeljanz, Jakvinus. The following terms were also used: economic analysis, economic assessment, economic evaluation, costeffectiveness, cost efficacy, and cost-utility. A detailed search strategy is available as electronic supplementary material. The CEA Registry was searched using the term 'ulcerative colitis'. In the CRD database, we omitted terms connected with the type of analysis.

The population included was described by the Population, Intervention, Comparator, Outcome (PICO) scheme: P—patients with moderate-to-severe UC (studies conducted in subpopulations were also included); I—biologic therapy with infliximab, adalimumab, golimumab, vedolizumab, etrolizumab or tofacitinib; C—not applicable, all other UC treatments were included; O—cost-effectiveness or cost-utility analyses.

The systematic review was independently carried out by two researchers, based on the same search strategy. The results for those searches were then compared; for disagreements, collaborative reassessment of the review results was performed to reach a final consensus.

2.2 Study Selection

Papers written in English were included. Neither country nor date filters were used. Original studies including primary data on the cost-effectiveness of biologics in UC, as well as reviews that led to previously undiscovered efficiency assessments, were included. Conference abstracts and posters were excluded as they did not provide the details valid for our review. The following exclusion criteria were implemented: a different intervention studied (i.e. pharmacotherapy other than biologics, surgery), different population (Crohn disease, other inflammatory bowel diseases) and different endpoints (i.e. cost assessments, clinical analyses, budget impact analyses). In this review, we included articles that fulfilled the inclusion criteria, defined within the PICO scheme, and which did not meet the exclusion criteria. During the selection of records, we used the Preferred Reporting Items for Systematic Reviews and the Meta-Analyses (PRISMA) statement.

2.3 Extraction and Presentation of Data

Data were extracted from the articles eligible. We used the Cochrane Handbook for Systematic Reviews of Intervention [8] to determine the terms collected: the first author's name, population, interventions, comparators, year of study, study design (assumptions and methods of the main analysis, i.e. the type of economic analysis, outcomes, perspective, country, time horizon, discount rates, cost reference date, currency, type of sensitivity analysis), results (total and incremental costs, total and incremental outcomes, incremental cost-effectiveness ratio/incremental cost-utility ratio [ICER/ICUR] value, conclusions on cost-effectiveness, results of sensitivity analysis). Two contributors discussed the discrepancies until a consensus was reached.

2.4 Quality Assessment

The reporting quality of the identified studies was assessed using the International Society of Pharmacoeconomics and Outcomes Research (ISPOR) Consolidated Health Economic Evaluation Reporting Standards (CHEERS) statement checklist. It does not evaluate the quality of conduct, however its objective is to ensure that the analyses are fully reported. This instrument consists of a 24-item checklist [9] verifying the presence of specific issues (e.g. perspective, comparators, time horizon) in the considered papers.

Two authors reviewed the studies and scored each question 0 (this particular aspect is not present) or 1 (this particular aspect is present). Any discrepancies were resolved by consensus.

2.5 Synthesis of Data

The identified studies were grouped depending on the type of biologics they investigated: infliximab, adalimumab, golimumab, vedolizumab, and other biologics. Cost-effectiveness was assessed based on conclusions presented by authors of the eligible studies.

3 Results

3.1 Study Selection

The systematic search identified 576 references. After removing duplicates, 489 remaining records were submitted for further selection based on the review of their titles and abstracts, when necessary. In the next step, we excluded 457 records for the following reasons: a different type of analysis (e.g. cost comparisons, budget impact analyses, clinical effectiveness analyses), secondary data (reviews), different intervention, different population (e.g. inflammatory bowel diseases in general, Crohn disease). Of the 32 records selected by full-text review, 16 were excluded due to secondary data (systematic reviews), ineligible publication type (conference abstracts), and different population or endpoints. The remaining 16 records were included in the review (Fig. 1). In our review, we found four reports for Canada, three for Poland, and two for England and Wales, as well as four for the UK. Additional evaluations were performed for The Netherlands, Iran and the US.

3.1.1 Infliximab

Most of the identified studies related to infliximab (75%; n = 12), two of which were cost-effectiveness analyses with response and/or remission and/or mucosal healing as

the outcomes [10, 11], and ten were cost-utility evaluations with quality-adjusted life-year (QALY) as an outcome. Four analyses used decision trees as an analytic technique [11–14], whereas the Markov modelling was used in the remaining evaluations. In the study by Wilson et al. [14], both the above modelling techniques were used (Table 1). Two analyses (Punekar and Hawkins [13] and Tsai et al. [15]) showed that infliximab is a cost-effective treatment option compared with standard care alone, whereas six other analyses revealed opposite conclusions [16–21], namely, the ICER/ICUR value exceeded the threshold

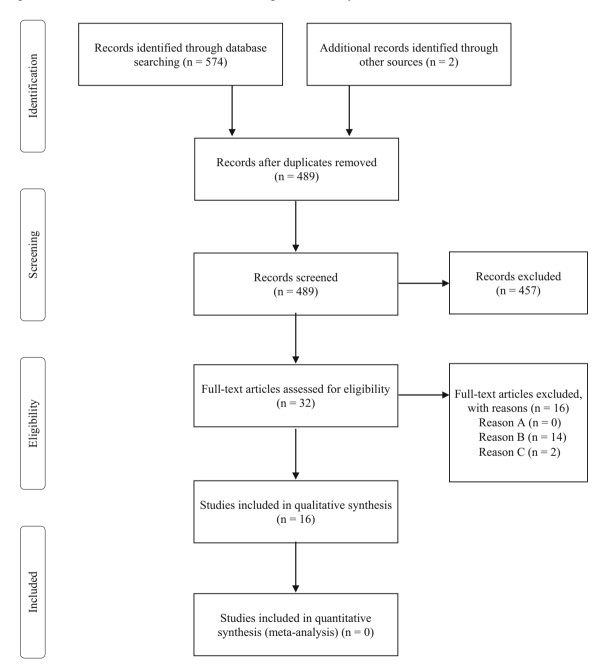


Fig. 1 PRISMA flow diagram showing the record selection process. Reason A—different intervention; Reason B—different type of study/ endpoint; Reason C—different population. *PRISMA* Preferred Reporting Items for Systematic Reviews and Meta-Analyses

Table 1 Characteristics of the included studies

Study, year	Population	Treatment	Study characteristics
Beilman	Moderate-to-severe active UC,	Adalimumab 160/80/	Type: Cost-utility analysis
et al., 2016 [23]	corticosteroid-dependent and/or did not respond to thiopurine therapy	40 mg, standard care	Model: Markov
			Horizon: 10 years
			Outcomes: QALY
			Discounting (costs/effects): 5%/5%
			Country: Canada
			Perspective: NA
			Reference year: NA
			Sponsor: Supported by the Centre of Excellence for Gastrointestinal Inflammation and Immunity Research
Chaudhary	Sever UC adult patients hospitalized	Infliximab 5 mg/kg,	Type: Cost-utility analysis
and Fan,	with an acute exacerbation of the	cyclosporine 2 mg/kg,	Model: Decision tree
2013 [12]	disease	surgical intervention	Horizon: 1 year
			Outcomes: QALY
			Discounting (costs/effects): 4%/1.5%
			Country: The Netherlands
			Perspective: National payer
			Reference year: 2010
			Sponsor: Merck & Co., Inc.
Aoradi et al.,	Moderate-to-severe UC	Infliximab 5 mg/kg,	Type: Cost-utility analysis
2016 [19]		conventional treatments	Model: Markov
			Horizon: 5 years
			Outcomes: QALY
			Discounting (costs/effects): 5%/NA
			Country: Iran
			Perspective: Public payer
			Reference year: 2014
			Sponsor: No
Punekar and Hawkins,	Moderate-to-severe UC patients hospitalized with an acute exacerbation of the disease	Infliximab 5 mg/kg, cyclosporine 4 mg/kg, surgery, standard care	Type: Cost-utility analysis
			Model: Decision tree
2010 [13]			Horizon: 1 year
			Outcomes: QALY
			Discounting (costs/effects): 3.5%/3.5%
			Country: England, Wales
			Perspective: Public payer
			Reference year: 2006–2007
			Sponsor: Schering-Plough Ltd
tawowczyk	Moderate-to-severe UC	Adalimumab 160/80/	Type: Cost-utility analysis
et al., 2016 [22]		40 mg, standard care	Model: Markov
			Horizon: 30 years
			Outcomes: QALY
			Discounting (costs/effects): 5%/3.5%
			Country: Poland
			Perspective: Public payer, social
			Reference year: 2015
			Sponsor: No

Table 1 continued

Study, year	Population	Treatment	Study characteristics
Stawowczyk	Moderate-to-severe UC	Golimumab 200/100/	Type: Cost-utility analysis
et al., 2016 [24]		50 mg, standard care	Model: Markov
			Horizon: 30 years
			Outcomes: QALY
			Discounting (costs/effects): 5%/3.5%
			Country: Poland
			Perspective: Public payer, social
			Reference year: 2015
			Sponsor: No
Stawowczyk	Moderate-to-severe UC	Infliximab 5 mg/kg,	Type: Cost-utility analysis
et al., 2016		standard care	Model: Markov
[21]			Horizon: 30 years
			Outcomes: QALY
			Discounting (costs/effects): 5%/3.5%
			Country: Poland
			Perspective: Public payer
			Reference year: 2015
			Sponsor: No
Fappenden	Moderate-to-severe UC in patients for	Infliximab 5 mg/kg,	Type: Cost-utility analysis
et al., 2016	whom at least one prior therapy has	adalimumab 160/80/	Model: Markov
[17]	failed	40 mg, golimumab 200 g/	Horizon: 60 years
		100/100 mg (50 mg), conventional non-biologic	Outcomes: QALY
		therapy, elective surgery	Discounting (costs/effects): 3.5%/3.5%
			Country: UK
			Perspective: Public payer, Personal Social Services
			Reference year: 2013/2014
			Sponsor: National Institute for Health Research Health Technology Assessment Programme
Foor et al.,	Moderate-to-severe UC	Infliximab 5 mg/kg, adalimumab 40 mg,	Type: Cost-effectiveness analysis
2015 [10]			Model: Markov
		golimumab 50/100 mg,	Horizon: 1 year
		conventional therapy	Outcomes: Remission, response
			Discounting (costs/effects): –
			Country: Canada
			Perspective: Public payer
			Reference year: 2013
			-
fsai et al	Moderate-to-severe UC	Infliximah 5 mg/kg	-
2008 [15]		standard care	
			-
			_
			-
[sai et al., 2008 [15]	Moderate-to-severe UC	Infliximab 5 mg/kg, standard care	Sponsor: Janssen Inc. Canada Type: Cost-utility analysis Model: Markov Horizon: 10 years Outcomes: QALY Discounting (costs/effects): 3.5%/3.5% Country: England, Wales Perspective: Public payer Reference year: 2006/2007 Sponsor: Schering-Plough Ltd

Table 1 continued

Study, year	Population	Treatment	Study characteristics
Ung et al.,	Moderate-to-severe UC	Infliximab 5 mg/kg,	Type: Cost-utility analysis
2014 [20]		standard care	Model: Markov
			Horizon: 10 years
			Outcomes: QALY
			Discounting (costs/effects): 5%/5%
			Country: Canada
			Perspective: Health system
			Reference year: 2013
			Sponsor: Supported by the Centre of Excellence for Gastrointestinal Inflammation and Immunity Research and the Alberta Innovates
Wilson et al.,	Moderate-to-severe UC	Vedolizumab 300 mg,	Type: Cost-utility analysis
2017 [14]		infliximab 5 mg/kg,	Model: Decision tree/Markov
		adalimumab 160/80/	Horizon: Lifetime
		40 mg, golimumab 100/50/50 mg,	Outcomes: QALY
		conventional therapy	Discounting (costs/effects): 3.5%/3.5%
			Country: UK
			Perspective: Public payer, Personal Social Services
			Reference year: 2012/2013
			Sponsor: Takeda Pharmaceuticals, AG
Xie et al.,	Moderate-to-severe refractory UC	Infliximab 5/10 mg,	Type: Cost-utility analysis
2009 [18]		adalimumab 160/80/	Model: Markov
		40 mg, usual care	Horizon: 5 years
			Outcomes: QALY
			Discounting (costs/effects): 5%/5%
			Country: Canada
			Perspective: Public payer
			Reference year: 2008
			Sponsor: No
Yokomizo	Moderate-to-severe UC	Infliximab 5/10 mg,	Type: Cost-effectiveness analysis
et al., 2016	Moderate-to-severe UC	adalimumab 160/80/	Model: Decision tree
[11]		40 mg, vedolizumab 300 mg	Horizon: 1 year
			Outcomes: Remission, mucosal healing
			Discounting (costs/effects): –
			Country: US
			-
			Perspective: Third party's
			Reference year: 2014 Sponsor: National Institute of Diabetes and Digestive
			and Kidney Diseases
Archer et al., 2016 [16]	Moderate-to-severe UC, after the failure of conventional treatment	Infliximab 5 mg/kg,	Type: Cost-utility analysis
		adalimumab 160/80/ 40 mg, golimumab	Model: Markov
		200/100/100 mg (50 mg),	Horizon: Lifetime
		conventional therapy,	Outcomes: QALY
		colectomy	Discounting (costs/effects): 3.5%/3.5%
			Country: UK
			Perspective: Public payer, Personal Social Services
			Reference year: 2013
			Sponsor: National Institute for Health Research

Table 1 continued

Study, year	Population	Treatment	Study characteristics
Essat et al., 2016 [25] ^a	Moderate-to-severe UC	Vedolizumab, conventional therapy, surgery, infliximab, adalimumab, golimumab	Type: Cost-utility analysis Model: Markov, decision-tree Horizon: Lifetime Outcomes: QALY Discounting (costs/effects): NA Country: UK Perspective: Public payer Reference year: NA Sponsor: Evidence Review Group

UC ulcerative colitis, QALY quality-adjusted life-years, NA not available, ERG Evidence Review Group

^aMethods and results of the ERG's own calculations were included

value in a particular country. In the study by Chaudhary and Fan [12], infliximab was shown to be cost-effective when compared with cyclosporine therapy and surgical intervention. Toor et al. [10] concluded that infliximab had the highest clinical efficacy (presented as response and remission), but also generated high costs compared with adalimumab and golimumab. In comparison with adalimumab and vedolizumab, infliximab turned out to be the most cost-effective treatment option when remission and mucosal healing were taken into account [11]. In the study by Archer et al. [16], infliximab was expected to be dominated by adalimumab (Table 2).

3.1.2 Adalimumab

Adalimumab was assessed in eight studies, six of which were cost-utility analyses displaying QALY as an outcome [14, 16, 17, 18, 22, 23]. The Markov modelling was used in seven studies, decision trees in two studies [11, 14], and, in the case of Wilson et al. [14], both analytic techniques (Markov modelling and the decision tree) were used (Table 1). Four studies showed that adalimumab is not a cost-effective treatment option when compared with standard care alone [16–18, 22], and only one study reported the opposite conclusion [23]. Toor et al. [10] concluded that adalimumab produced higher cost per remission and cost per response, when compared with standard care, than other biologics, i.e. infliximab and golimumab. In the study by Yokomizo et al. [11], adalimumab was dominated (proving less effective and more costly) by infliximab, when mucosal healing was included as an outcome (Table 2).

3.1.3 Golimumab

Golimumab was included as a treatment option in UC in five studies, four of which were cost-utility analyses [14, 16, 17, 24], and one included remission and response

as the outcomes [10]. All the studies used the Markov modelling and one study additionally used a decision tree [14] (Table 1). Stawowczyk et al. [24], Tappenden et al. [17] and Archer et al. [16] concluded that golimumab is not a cost-effective treatment option when compared with standard care alone. In the study by Toor et al. [10], golimumab had the lowest cost of 1 additional year of remission and response compared with standard care, among other biologics included, i.e. infliximab and adalimumab. Wilson et al. [14] showed that golimumab was dominated by vedolizumab (Table 2).

3.1.4 Vedolizumab

Only three studies assessed the cost-effectiveness of vedolizumab in the treatment of UC; one used a decision tree [11] and two used both the decision tree and Markov modelling [14, 25]. Wilson et al. [14] compared vedolizumab with other biologics used in UC, i.e. infliximab, adalimumab and golimumab, and concluded that it is a cost-effective treatment option (Table 1). Vedolizumab generated the highest outcomes (QALYs) and was cheaper than infliximab and golimumab (therefore it dominated both comparators). Only adalimumab proved to be cheaper than vedolizumab, but also less effective-the ICER for vedolizumab compared with adalimumab did not exceed the threshold value. Yokomizo et al. [11] included mucosal healing as an outcome and showed that vedolizumab was dominated by infliximab. Essat et al. [25] described the results of Evidence Review Group (ERG) assessment of vedolizumab. Based on the ERG's own calculations, it was shown that vedolizumab is expected to be dominated by surgery (Table 2).

3.2 Quality Assessment

The results of quality assessment with the ISPOR CHEERS statement checklist are shown in Table 3. The study by

Table 2 Results of the included studies

Study, year	Total costs	Total outcomes	ICER	Authors' conclusions
Beilman et al., 2016 [23]	SC = US\$97,000 ADA = US\$107,000	SC = 3.154 ADA = 3.321	Per QALYG: ADA vs. SC = US\$59,000	ADA is cost-effective compared with SC
Chaudhary and Fan, 2013 [12]	IFX = ϵ 17,062 CSP = ϵ 14,784 S = ϵ 13,979	IFX = 0.80 CSP = 0.70 S = 0.58	Per QALYG: IFX vs. CSP = \notin 24,277 IFX vs. S = \notin 14,639	IFX is cost-effective compared with CSP and S
Moradi et al., 2016 [<mark>19</mark>]	IFX = US\$77,138 $SC = US$985$	IFX = 3.56 SC = 3.24	Per QALYG: IFX vs. SC = US\$240,903	IFX is not cost-effective compared with SC
Punekar and Hawkins, 2010 [13]	$S = \pounds 17,067$ $CSP = \pounds 18,122$ $SC = \pounds 18,524$ $IFX = \pounds 19,847$	S = 0.58 CSP = 0.70 SC = 0.68 IFX = 0.80	Per QALYG: IFX vs. $CSP = \pounds 19,545$	IFX is cost-effective compared with CSP, S, SC
Stawowczyk et al., 2016 [22]	Public payer: $ADA = \epsilon 20,598$ $SC = \epsilon 9950$ Social: $ADA = \epsilon 93,765$ $SC = \epsilon 83,770$	ADA = 15.204 SC = 15.064	Per QALYG: Public payer: ADA vs. $SC = \epsilon 76,120$ Social: ADA vs. $SC = \epsilon 71,457$	ADA is not cost-effective compared with SC
Stawowczyk et al., 2016 [24]	Public payer: GOL = PLN93,321 SC = PLN45,502 Social: GOL = PLN302,848 SC = PLN257,092	GOL = 19.241 SC = 19.118	Per QALYG: Public payer: GOL vs. SC = 391,252 PLN Social: GOL vs. SC = 374,377 PLN	GOL is not cost-effective compared with SC
Stawowczyk et al., 2016 [21]	IFX = PLN99,522 $SC = PLN29,642$	IFX = 14.296 SC = 14.123	Per QALYG: IFX vs. SC = PLN402,420	IFX is not cost-effective compared with SC
Tappenden et al., 2016 [17]	$S = \pounds 56,268$ ADA = \pounds 91,222 IFX = \pounds 96,595 GOL = \pounds 90,087 SC = \pounds 73,620	S = 14.71 ADA = 10.82 IFX = 10.81 GOL = 10.63 SC = 10.47	Per QALYG: S vs. ADA/IFX/GOL/SC–S dominated all therapies ADA vs. SC = \pounds 50,278 IFX vs. SC = \pounds 67,574 ^a GOL vs. SC = \pounds 102,919 ^a	Colectomy dominated all medical treatment. ADA, IFX ^a , GOL ^a are not cost- effective compared with SC
Toor et al., 2015 [10]	NA	NA	Per remission: GOL 50 vs. SC = CAN $$1048$ GOL 100 vs. SC = CAN $$935$ IFX vs. SC = CAN $$1975$ ADA vs. SC = CAN $$7430$ IFX vs. GOL 100 = CAN $$14,659$ ADA vs. GOL 100 = - CAN $$3324$ Per response:	GOL 100 and GOL 50 have the lowest cost of additional 1 year of remission and response; IFX has the highest efficacy, but also high costs, ADA produced the highest cost/remission and response

GOL 50 vs. SC = CAN\$770 GOL 100 vs. SC = CAN\$701 IFX vs. SC = CAN\$1311 ADA vs. SC = CAN\$2361 IFX vs. GOL 100 = CAN\$4753 ADA vs. GOL 100 = - CAN\$4019

Table 2 continued

Study, year	Total costs	Total outcomes	ICER	Authors' conclusions
Tsai et al., 2008 [15]	Responders only: IFX = $\pounds 66,460$ SC = $\pounds 45,798$ Remission only: IFX = $\pounds 53,874$ SC = $\pounds 46,259$	Responders only: IFX = 4.591 SC = 3.838 Remission only: IFX = 4.154 SC = 3.767	Per QALYG: Responders only: IFX vs. $SC = \pounds 27,424$ Remission only: IFX vs. $SC = \pounds 19,696$	IFX is cost-effective compared with SC
Ung et al., 2014 ^b [20]	SC = US\$86,000 IFX = US\$98,000	SC = 3.707 SC = 3.204 IFX = 3.284	Per QALYG: IFX vs. SC = US\$152,000	IFX is not cost-effective compared with SC
Wilson et al., 2017 [14]	$VED = \pounds 199,431$ IFX = \\pounds 206,066 ADA = \\pounds 194,765 GOL = \\pounds 200,018	VED = 14.077 IFX = 13.788 ADA = 13.872 GOL = 13.809	Per QALYG: VED vs. IFX—VED dominated IFX VED vs. ADA = £22,775 VED vs. GOL—VED dominated GOL	VED is cost-effective compared with IFX, ADA, GOL
Xie et al., 2009 [18]	SC = CAN\$24,268 IFX ADA = CAN\$82,756 IFX 5 IFX 10 ADA = CAN\$101,272	SC = 2.015 IFX ADA = 2.178 IFX 5 IFX 10 ADA = 2.149	Per QALYG: IFX ADA vs. $SC = CAN$ \$358,088 IFX 5 IFX 10 ADA vs. SC = CAN\$575,540	Anti-TNFα therapies are not cost-effective compared with SC
Yokomizo et al., 2016 [11]	IFX $5 = US$99,171$ IFX $10 = US$123,653$ ADA = US\$316,378 VED = US\$301,969	NA	Per mucosal healing: IFX 5 vs. IFX 10 = US\$1,243,310 IFX 5 vs. ADA—IFX dominated ADA IFX 5 vs. VED—IFX dominated VED	IFX is the most cost-effective treatment option
Archer et al., 2016 [16]	$S = \pounds 56,267.73$ ADA = \pounds 91,221.71 IFX = \pounds 96,594.62 GOL = \pounds 90,086.69 SC = \pounds 73,619.77	S = 14.71 ADA = 10.82 IFX = 10.81 GOL = 10.63 SC = 10.47	IFX vs. ADA—ADA dominated IFX ADA vs. SC = \pounds 50,278 IFX vs. SC = \pounds 67,573 ^a GOL vs. SC = \pounds 102,918 ^a	Surgery dominated all other options. ADA dominated IFX
Essat et al., 2016 [25] ^c	NA	NA	Surgery is an option: VED vs. S—S dominated VED Surgery is not an option, mixed ITT population: VED vs. SC = £53,084 Surgery is not an option, anti-TNFα failure population: VED vs. SC = £48,205	When surgery is an option it dominated VED. When surgery is not an option, VED was not cost-effective compared with SC

CAN\$1 = US\$0.794; €1 = US\$1.184; £1 = US\$1.324; 1 PLN = US\$0.278

IFX infliximab, *ADA* adalimumab, *GOL* golimumab, *GOL* 50 golimumab 50 mg, *GOL* 100 golimumab 100 mg, *VED* vedolizumab, *SC* standard/ usual care/conventional treatment, *S* surgery, *CSP* cyclosporine, *QALYG* quality-adjusted life-years gained, *ICER* incremental cost-effectiveness ratio, *NA* not available, *TNF* tumor necrosis factor

^aOur own calculations based on available data

^bResults based on data from clinical studies

^cResults of the ERG's own calculations were included

Essat et al. [25] was not assessed as it presents the results of a Health Technology Assessment (HTA) rather than typical economic analysis.

The identified economic analyses obtained a rather high average result of the ISPOR CHEERS statement checklist, equalling 21 points, with a minimum and maximum value of 14 and 23, respectively. Only one study obtained the minimum value [19], whereas the maximum value was reached by three studies [14, 22, 24]. The most common value was the average value, i.e. 21 points (see electronic supplementary material).

All the included studies had adequate titles, which identified the study as an economic evaluation (the most common phrases were cost-effectiveness, cost-utility, economic evaluation) and described the compared interventions. Studies failed to report relevant aspects on the system in which the decision needed to be made—adequate information was presented in only six studies (40%)

Table 3 The results of quality assessment of included studies with ISPOR CHEERS statement checklist

Section/item	Item no.	Recommendation	Number of studies in line with recommendation	Percentage of studies in line with recommendation
Title and abstract				
Title	1	Identify the study as an economic evaluation, or use more specific terms such as "cost-effectiveness analysis" and describe the interventions compared	15	100.0
Abstract	2	Provide a structured summary of objectives, perspective, setting, methods (including study design and inputs), results (including base-case and uncertainty analyses), and conclusions	15	100.0
Introduction				
Background and objectives	3	Provide an explicit statement of the broader context for the study. Present the study question and its relevance for health policy or practice decisions	15	100.0
Methods				
Target population and subgroups	4	Describe characteristics of the base-case population and subgroups analyzed including why they were chosen	15	100.0
Setting and location	5	State relevant aspects of the system(s) in which the decision(s) need(s) to be made	6	40.0
Study perspective	6	Describe the perspective of the study and relate this to the costs being evaluated	14	93.3
Comparators	7	Describe the interventions or strategies being compared and state why they were chosen	15	100.0
Time horizon	8	State the time horizon(s) over which costs and consequences are being evaluated and say why appropriate	15	100.0
Discount rate	9	Report the choice of discount rate(s) used for costs and outcomes and say why appropriate	13	86.7
Choice of health outcomes	10	Describe what outcomes were used as the measure(s) of benefit in the evaluation and their relevance for the type of analysis performed	15	100.0
Measurement of effectiveness	11a	Single study-based estimates: Describe fully the design features of the single effectiveness study and why the single study was a sufficient source of clinical effectiveness data	14	93.3
	11b	Synthesis-based estimates: Describe fully the methods used for the identification of included studies and synthesis of clinical effectiveness data		
Measurement and valuation of preference- based outcomes	12	If applicable, describe the population and methods used to elicit preferences for outcomes	13	86.7
Estimating resources and costs	13a	Single study-based economic evaluation: Describe approaches used to estimate resource use associated with the alternative interventions. Describe primary or secondary research methods for valuing each resource item in terms of its unit cost. Describe any adjustments made to approximate to opportunity costs	15	100.0
	13b	Model-based economic evaluation: Describe approaches and data sources used to estimate resource use associated with model health states. Describe primary or secondary research methods for valuing each resource item in terms of its unit cost. Describe any adjustments made to approximate to opportunity costs		

Table 3 continued

Section/item	Item no.	Recommendation	Number of studies in line with recommendation	Percentage of studies in line with recommendation
Currency, price date, and conversion			13	86.7
Choice of model	-		14	93.3
Assumptions	16	Describe all structural or other assumptions underpinning the decision-analytic model	13	86.7
Analytic methods	17	Describe all analytic methods supporting the evaluation. This could include methods for dealing with skewed, missing, or censored data; extrapolation methods; methods for pooling data; approaches to validate or make adjustments (e.g., half-cycle corrections) to a model; and methods for handling population heterogeneity and uncertainty	12	80.0
Results				
Study parameters	18	Report the values, ranges, references, and if used, probability distributions for all parameters. Report reasons or sources for distributions used to represent uncertainty where appropriate. Providing a table to show the input values is strongly recommended	14	93.3
Incremental costs and outcomes	19	For each intervention, report mean values for the main categories of estimated costs and outcomes of interest, as well as mean differences between the comparator groups. If applicable, report incremental cost-effectiveness ratios	15	100.0
Characterizing uncertainty	20a	Single study-based economic evaluation: Describe the effects of sampling uncertainty for estimated incremental cost, incremental effectiveness, and incremental cost-effectiveness, together with the impact of methodological assumptions (such as discount rate, study perspective)		
	20b	Model-based economic evaluation: Describe the effects on the results of uncertainty for all input parameters, and uncertainty related to the structure of the model and assumptions	14	93.3
Characterizing heterogeneity	21	If applicable, report differences in costs, outcomes, or cost- effectiveness that can be explained by variations between subgroups of patients with different baseline characteristics or other observed variability in effects that are not reducible by more information	0	0.0
Discussion				
Study findings, limitations, generalizability, and current knowledge	22	Summarize key study findings and describe how they support the conclusions reached. Discuss limitations and the generalizability of the findings and how the findings fit with current knowledge	13	86.7
Other				
Source of funding	23	Describe how the study was funded and the role of the funder in the identification, design, conduct, and reporting of the analysis. Describe other nonmonetary sources of support	13	86.7
Conflicts of interest	24	Describe any potential for conflict of interest among study contributors in accordance with journal policy. In the absence of a journal policy, we recommend authors comply with International Committee of Medical Journal Editors' recommendations	12	80.0

[14, 16, 18, 21, 22, 24]. The healthcare system and reimbursement status of the drugs analysed in the particular countries were rarely described. Study perspective was rather properly described, only in one study was the information missing [23]. In a majority of revealed analyses [2, 12-19, 21, 22] a public payer perspective was applied, and in the case of two evaluations [22, 24] a social point of view was employed. One study [11] did not report information on discount rates for costs and outcomes, but the time horizon of this study was short (1 year) and discounting was not needed. In another study [19], there was no information regarding the discount rate for outcomes, and only the rate for costs was reported. Preferences for outcomes (utility weights) were included and properly described in 13 studies (87%), whereas in the studies by Toor et al. [10] and Yokomizo et al. [11] they were not included because they employed a different type of analvsis (cost-effectiveness, with remission and response as outcomes). In some studies, information on the currency and/or exchange rates was missing. Beilman et al. [23] did not describe the year of reported costs, and Xie et al. [18] presented 2008 costs in Canadian dollars but did not present the exchange rate.

Only one study [10] did not reveal the values for main study parameters; the authors presented only the cost per remission and cost per response for biologics versus conventional therapy and other biologics. The heterogeneity was not assessed in any of the identified studies as no study included subgroups.

3.3 Drivers of Cost-Effectiveness

3.3.1 Uncertainty Within the Study

In 14 studies (88%), sensitivity analyses were carried out, and only in the study by Moradi et al. [19] were the effects of sampling uncertainty for estimated results not described as the authors failed to undertake the probabilistic sensitivity analysis. In the study by Essat et al. [25], where the results of HTA were presented, no information on sensitivity analysis was available. Both deterministic and probabilistic sensitivity analyses were performed, and one-way sensitivity analysis was the most often used type of deterministic sensitivity analyses. The most common sensitive variables included in the analyses were the parameters of efficacy (remission/response rates, transition probabilities), utility weights, discount rates, cost parameters, and patients' characteristics (weight). The majority of studies revealed that the change of utility weights [12, 13, 16-18, 20-24] and clinical parameters, i.e. remission/response rates, transition probabilities, [13, 14, 17, 18, 21, 22, 24] had the highest influence on analysis results (ICER/ICUR values). The change of patients' characteristics. i.e. body weight, [12, 13] also had a significant influence on the results in some cases. The costs were less often included in the sensitivity analysis; however, they had a significant influence on the results in a few cases, i.e. the cost of response to adalimumab treatment [23], infliximab cost [20], health-state costs [14], and drug costs [11]. In two studies, the change of the duration of treatment had a great influence on the results of the analysis [10, 24]. The variable related to surgery, i.e. the probability of surgical complications, was the least sensitive [20, 21, 23, 24].

3.3.2 Uncertainty Between Studies

The results of some studies were also compared to estimate the impact of model assumptions on the results of the analysis. We chose analyses with the same outcomes (i.e. QALY), the same comparator (i.e. standard care) and the same intervention (i.e. infliximab, as the highest number of studies related to this biologic). Seven studies met the above criteria [15-21], but two studies had to be subsequently excluded as one included different treatment strategies (infliximab 5 mg/kg + infliximab 10 mg/kg + adalimumab,and infliximab + adalimumab [18]), and one provided results separately for responders and patients in remission [15]. The range of the ICUR value in the remaining studies ranged from US\$89,468 [16, 17] to US\$240,903 [19]; however, the conclusions relating to cost-effectiveness were the same: infliximab was not a cost-effective option in the treatment of UC compared with standard care alone. All analyses used the Markov modelling, but the time horizon differed significantly from 5 years [19] to lifetime [16], and the lowest and highest ICUR values were obtained for analyses with the longest and shortest time horizon, respectively. The other model assumptions were similar-discount rates: 5% in the majority of cases, or 3.5% in the studies by Tappenden et al. [17], Archer et al. [16] and Stawowczyk et al. [21] (only for effects); study perspective: public payer/health system; reference year: 2013-2015; utility weights: based on the studies by Arseneau et al. [26] (all studies) and Woehl et al. [27] (studies [16, 17, 21]).

3.4 Model Structure

To compare the structures of the models, we first chose analyses that used the Markov modelling as this was the methodology most often performed; we identified 12 such analyses [14–25], but in study by Essat et al. [25] the model structure was not presented in a form of diagram and no

detailed information was provided. That was the reason why this study was excluded.

Beilman et al. [23] and Ung et al. [20] compared two scenarios—when adalimumab is available and when it is not available. Several health states were identified: unwell during treatment, response (to pharmacotherapy or surgery), complications (after pharmacotherapy or surgery) and death. Chronic pouchitis was analysed regardless of complications after surgery and was considered as a separate state.

The structure of the model presented by Moradi et al. [19] was less sophisticated and included only three states, i.e. remission, response and surgery.

All three models used in the studies by Stawowczyk et al. [21, 22, 24] were similar and included the following states: remission (after pharmacotherapy or surgery), response, active disease, surgery, and complications after surgery. The model for infliximab [21] additionally included the second-line treatment with adalimumab (after failure of infliximab treatment).

Similar states and model structures were included in the study by Tsai et al. [15], Archer et al. [16] and Tappenden et al. [17]; however, only first-line biologic treatment was included. Tsai et al. [15] included the following health states: temporary discontinuers, remission, mild, moder-ate-severe, surgery, post-surgery remission and post-surgery complications. Archer et al. [16] and Tappenden et al. [17] divided the model structure into four parts, i.e. on biological treatment, non-biological treatment, post-surgery, and dead. The first and second parts included the same states: active UC, response, and remission, while the third and fourth parts included only one state: post-surgery (with/without complications) and dead, respectively.

Wilson et al. [14] did not include the response state, but instead two others were included: mild with a Mayo score of 3-5 and moderate–severe with a Mayo score of 6-12. Additionally, six other states were included: remission with a Mayo score of 0-2, surgery (from moderate–severe state), post-surgery remission, post-surgery complications, discontinue and death.

In the analysis by Xie et al. [18], three strategies were included: usual care, infliximab 5 mg/kg followed by adalimumab, and infliximab 5 mg/kg followed by infliximab 10 mg/kg and then adalimumab. The states in the model were similar to those in other studies, i.e. response, remission, surgery, complications, and active disease.

We also compared the model structures in analyses where the decision trees were the chosen methods of modelling [11-14]; a significant heterogeneity in the approaches used in the considered analyses was revealed. The structure of the decision trees and the assumptions included in the modelling varied among the studies.

Chaudhary and Fan [12] used the decision tree to compare infliximab, cyclosporine and surgery over a 1-year

time horizon. The following states were included: surgery or remission after pharmacotherapy, and remission or complication after surgery.

In another study, infliximab treatment was compared with standard care, cyclosporine and surgery [13]. The states included initial remission and early surgery, and initial remission transitioned to ongoing remission and late surgery states.

Quite a simple version of the decision tree was used in the study by Wilson et al. [14] for the evaluation of biologic induction. Patients who responded to therapy during the induction period and did not discontinue as a result of adverse effects, remained on maintenance treatment (entered Markov model). Patients who did not respond during the induction phase, lost response during the maintenance phase, or discontinued due to adverse events switched to induction with conventional therapy [14].

No detailed structure of the decision analytical model was provided in the case of the decision tree used to compare the cost-effectiveness of the first-line treatment with adalimumab, infliximab or vedolizumab to induce mucosal healing [11]. The authors presented only a simplified scheme, without particular states or transitions. The lack of data on the methods used makes it impossible to compare the model structure with other analyses.

3.5 Data Sources

The half of analyses included the results of single clinical trials collected in the review of literature in order to estithe transition probabilities in the mate model [10–12, 15, 19, 22–24]. In other studies, the meta-analyses and indirect comparisons (network meta-analyses) of the identified clinical trials were conducted to obtain the clinical data used as input for simulations [13, 14, 16–18, 20, 21, 25].

The country databases were a source of information to estimate the unit costs of drugs and medical services in 13 of 16 identified analyses [10–19, 21, 22, 24]. In only two studies [20, 23] was the literature search conducted to obtain the cost data. In study by Essat et al., no information on cost sources was provided [25].

4 Discussion

We identified 16 economic analyses, mainly cost-utility analyses, which assessed the cost-effectiveness of biologic drugs (infliximab, adalimumab, golimumab, and vedolizumab) in the treatment of UC. The assessed interventions were mainly compared with standard care alone, but cyclosporine, surgical intervention and other biologics were also included as comparators in some cases. The highest number of economic analyses related to infliximab, and only three of the analyses included vedolizumab, including one HTA report. The vast majority of studies used Markov modelling and only five studies used the decision trees. All Markov models included response, remission, and surgery states, but response state was, in some cases, replaced by mild state. The second-line biologic treatment was rarely included in the analysis. The structure of decision trees varied significantly between the studies. The most commonly applied time horizon was lifetime (30 years or more), which was adopted in 44% of analyses (7 of 16), but a 1-year time horizon was also employed quite often (27% of the analyses). In our opinion, a lifetime horizon is more valuable for chronic diseases as the longer period of observation provide more sophisticated information on consequences of therapy compared with a 1-year horizon. The QALY was used as an outcome in 88% of the analyses, while the other analyses used remission, response and mucosal healing. Cost-effectiveness studies on new treatments should take into consideration the impact of the new interventions on patient productivity and a corresponding economic burden for the society. Therefore, it should be emphasized that only two of the economic analyses also carried out the calculations from a social perspective, and took into account the impact of treatment on work productivity, which seems quite unusual because UC, a condition usually diagnosed in middle-aged patients, generates a significant loss of productivity; therefore, indirect costs could even exceed direct healthcare costs. Both analyses were conducted in Polish settings [22, 24]. Further studies should be performed to measure the influence of productivity loss on the cost-effectiveness of biologic drugs in UC.

The quality of the included analyses was evaluated as high, based on review of methodologies used and total score of the CHEERS reporting quality. A majority of valuable and important information referring to reporting quality was provided in the reports collected, excluding the description of national healthcare systems or reimbursement status of assessed intervention, which was quite rarely presented in the analyses.

According to reimbursement criteria, the acceptable cost-effectiveness of biologics compared with standard care alone was revealed in only three studies. The performed sensitivity analyses demonstrated that the costeffectiveness results (ICER/ICUR values) were the most sensitive to the change of utility weights and clinical parameters (response/remission rates, transition probabilities). The between-study comparison may suggest that the longer the time horizon, the lower the ICUR values. This may result from the fact that UC is a lifelong disease and its effects revealed during the whole lifespan.

The included analyses were country-specific as they included the costs of therapies valid for each specified country. Clinical outcomes can be transferred to other countries and generalized; however, cost inputs are largely country-specific, which in turn limits the transferability and generalizability of the results and conclusions to other countries. Another issue is the differences in healthcare systems and reimbursement policies between countries, and also the methods of inpatient or outpatient care, which may have a significant influence on the results and final conclusions of economic evaluations.

We critically reviewed the collected publications but also discussed the model structures, cost sources, model assumptions, model validation and key drivers for costeffectiveness analyses to provide a substantial contribution over the existing reviews in the topic.

We limited the final review to studies written in English, which may be considered as a limitation, as this approach produced the potential omission of relevant papers. Several identified analysis were conducted directly by manufacturers and their results were published in a form of an HTA report. We did not include the results of manufacturers' calculations, which may also be a limitation, but the results of analysis carried out by HTA agencies were taken into account. Such an approach was implemented because manufacturers' analysis, which are industry-sponsored evaluations, may be biased. Another limitation is that the quality assessment was not performed for one analysis [25]. The analysis results were presented as an ERG assessment and did not include the structure typical for economic evaluations. The heterogeneity of model assumptions was the reason why meta-analysis was not performed for identified studies, which also constitutes a limitation of this review.

Despite the above limitations, we included the majority of cost-effectiveness studies currently available, which enabled us to evaluate the overall quality and identify key drivers of cost-effectiveness.

This study is the most comprehensive review that incorporates economic evaluations referring to UC treatments. It includes quality assessment with a validated instrument and in-depth analysis of cost-effectiveness key drivers.

There is a need for studies that will provide evidence for the effectiveness of long-term treatment of UC and resolve the uncertainty associated with the key drivers of efficiency.

5 Conclusions

We identified well-reported economic analyses providing conclusions on the cost-effectiveness of infliximab, adalimumab, golimumab and vedolizumab therapy in UC. Conclusions based on the performed cost-effectiveness analyses were not unequivocal. The key drivers identified in the review were utility weights and clinical inputs. **Data Availability Statement** We made every effort to provide a sophisticated and complex set of data while performing the review. Any data valid during the current review were presented in the electronic upplementary material.

Authors Contributions ES and PK contributed to the study rationale and design, and performed a systematic review and quality assessment of identified studies. ES carried out interpretation of the data. Both authors prepared the draft of the manuscript, contributed to editing and revising the manuscript for important intellectual contributions, and approved the final version submitted for publication.

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Compliance with Ethical Standards

Conflict of interest Ewa Stawowczyk and Paweł Kawalec declare that they have no conflicts of interest.

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