

1 currently used in inflammatory arthritis across age, and will examine any significant
2 differences between ADA prevalence, titres and timing of development, as well as ADA impact
3 on therapeutic drug levels, clinical efficacy and side-effects between paediatric and adult
4 patients. In addition, this paper will investigate factors associated with differences in
5 immunogenicity across biologic agents used in inflammatory arthritis, and their potential
6 therapeutic implications.

7

8 **Introduction**

9 The discovery and clinical use of biologic treatments in the management of inflammatory
10 arthritis in children and adults has been associated with significant clinical benefits as well as
11 advances in understanding the pathogenesis of different types of inflammatory arthritis.
12 Immunogenicity to biologic treatments is an unwanted immune reaction against a therapeutic
13 antigen. This immune reaction generates anti-drug-antibodies (ADA), which could counteract
14 the therapeutic effects of the biologic treatment and, in rare cases, induce adverse reactions (1,
15 2).

16 It has become increasingly recognised that biologic treatment duration, mode, rate and route of
17 administration, and more specifically the type of biologic therapeutic (e.g. monoclonal
18 antibodies - mAbs versus recombinant fusion proteins) are all factors that influence the risk of
19 immunogenicity (3). In addition, individual patient factors, such as genetic background (4),
20 disease type (5), and concomitant use of disease modifying anti-rheumatic drugs (DMARDs)
21 (6), all contribute differentially to the formation of ADA. Recent research has been focused on
22 highlighting the genetic risk for developing ADA: e.g. HLA-DRB1*15 was associated with
23 increased the risk for developing high ADA levels to interferon (IFN) β -1a treatment in multiple
24 sclerosis, while HLA-DQA1*05 decreased this risk (7), and HLA-DQA1*05 was associated
25 with increased ADA prevalence across various biologics and autoimmune diseases (8). Other
26 factors such as smoking and infections are also associated with increased risk (8, 9), whereas
27 concomitant use of antibiotics and immunosuppressant medication are associated with
28 decreased immunogenicity risk (8). In addition, the manufacturing process of various biologic
29 agents, in particular their contamination with low-level host proteins, is a major contributor to
30 immunogenicity (10).

31 Therapeutic drug monitoring and immunogenicity testing comprise measurement of trough
32 drug levels and ADA. The most widely used ADA detection methods are bridging ELISA

1 (which use labelled therapeutic mAbs) and radioimmunoassay (RIA), while other new methods
2 such as competitive displacement and tandem mass spectrometry have also been proposed (11).
3 Currently, most mAbs on the market are humanised or fully human; however, they still carry
4 immunogenic risk. This could be attributed to anti-idiotypic reactivity, which is a common
5 reaction of the immune system to the appearance of any novel antibody (12).

6 The molecular mechanisms leading to generation of ADA are not completely elucidated and a
7 detailed discussion of immune mechanisms is beyond the scope of this review (for a recent
8 review see (13)). One basis for ADA generation involves the capacity of the human immune
9 system to recognise “non-self”. Since the first therapeutic mAbs of murine origin were
10 developed, further efforts have now been made to improve their performance and decrease
11 their immunogenicity. The continuous advancement in recombinant DNA technologies has led
12 to the development of chimeric (fused human-murine mAbs) and humanised mAbs. Chimeric
13 antibodies were developed by replacing the constant region of murine mAbs with human
14 components and the humanised mAbs are constituted entirely of human sequences, with the
15 exception of the complementarity determining regions (CDRs) of the variable (V) regions
16 which are of mouse-sequence origin. Subsequently, the advanced antibody engineering
17 achieved the production of fully human antibodies where antigen specificity has been selected
18 either *in vivo* in genetically modified mice or by antibody engineering processes combined with
19 screening (14). Many factors contribute to differences in immunogenicity, from
20 biopharmaceutical properties related to downstream processing and drug formulation (15) to
21 patient individual characteristics, including the antigen burden which correlates with their
22 disease activity (16).

23

24 Both ELISAs and RIAs detect only free circulating ADAs; therefore, they can be associated
25 with false negative results in the context of presence of ADA-immune complexes which are
26 detectable only if they exceed in concentration the circulating drug levels (17, 18). In one study,
27 ELISA was more sensitive in detecting ADA when present in high titres than RIA, while in
28 patients with ADA detected by RIA but not by ELISA only the drug levels were significantly
29 associated with treatment response to adalimumab (19). Interestingly, measuring drug levels
30 and drug clearance alone has also been shown to be a reliable predictor for ADA in RA and
31 juvenile idiopathic arthritis (JIA) patients (20)(21). Several studies concluded that although
32 ADA were not independently associated with treatment response, they may be helpful in
33 determining the cause of low drug levels and guide therapeutic decisions (22, 23).

1

2 The presence of ADAs may be associated with reduced clinical efficacy through two main
3 mechanisms. ADA that compete with the cytokine binding site (the Fab fragment of the
4 therapeutic agent) have neutralising properties as they block the pharmacological function of
5 the drug. ADA directed against the Fc fragment (more frequently targeting the junction
6 between Fc and Fab) lead to formation of immune complexes associated with enhanced drug
7 clearance may also influence the clinical response to biologic treatment through leading to sub-
8 optimal (sub-therapeutic) drug levels (24). Therefore, based on their specificity ADA can be
9 grouped as neutralising (when they target the antigen binding sites of the therapeutic drug) or
10 non-neutralising (when they recognize epitopes away from the drug binding site, therefore not
11 directly impairing the efficacy of the drug)(3).

12 Here we review the evidence of impact of ADA against various biologic therapeutics used for
13 treatment of inflammatory arthritis in adults and children as there are no previous reports
14 investigating immunogenicity across age. This review focuses on depicting differences
15 between ADA prevalence, titres and timing of development, as well as impact on therapeutic
16 drug levels, clinical efficacy and side-effects in children compared to adults with inflammatory
17 arthritis. Where data is available, we will also investigate the clinical predictors for ADA
18 development, as well as the influence of additional DMARD therapy on ADA development
19 and biologic drug retention.

20

21 **Neutralising ADA against mAbs targeting TNF- α were more prevalent than ADA**
22 **against fusion proteins (etanercept and biosimilars) while the kinetic of ADA generation**
23 **varied across anti TNF- α agents in adult and paediatric inflammatory arthritis studies**

24 Many studies have reported the presence of ADA against anti-TNF- α inhibitors used to treat
25 different types of inflammatory arthritis including etanercept (fusion protein of the
26 extracellular ligand-binding portion of the human 75KD p75 TNF receptor (TNFR) linked to
27 the Fc portion of human IgG1), adalimumab (fully human mAb), certolizumab (humanised
28 antibody Fab' fragment), golimumab (human IgG1 κ monoclonal antibody) or infliximab (a
29 chimeric mAb) (Table 1). The general observation is that ADA against etanercept have a lower
30 prevalence compared to ADA against adalimumab or infliximab (25). Furthermore,
31 comparative studies show that ADA to human/humanised (adalimumab, certolizumab,

1 golimumab) and chimeric (infliximab) anti-TNF- α therapeutic mAbs are largely neutralising
2 (26), while the ADA against etanercept are predominantly non-neutralising (27).

3

4 In adults, the rates of ADA formation against infliximab range from 8-62% in rheumatoid
5 arthritis (RA), 15-33% for psoriatic arthritis (PsA) and 6.1-69% for ankylosing spondylitis
6 (AS) (28) (Table 1). ADA against infliximab have also been shown to be associated with lower
7 serum biologic drug concentrations in adult inflammatory arthritis patients (27-35). There is a
8 paucity of studies investigating the timing of development of ADA against various anti-TNF-
9 α agents: evidence suggests that longer exposure to infliximab increases immunogenicity; e.g.
10 ADA against infliximab in adults with RA occurred after the first 10 infusions (23.4 ± 2.4
11 weeks) , while ADA were detected in 25% of JIA patients after 52 weeks and in 37% at 204
12 weeks (36-38). The dose of biologic agent as well as patients' age could influence
13 immunogenicity: a higher incidence of ADAs was observed in patients treated with infliximab
14 3mg/kg (38%), compared to 6 mg/kg (12%) (37), while a significantly higher prevalence of
15 ADA was found in younger children (ADA positive mean age 7.01 years vs. ADA negative
16 9.88 years, $p = 0.003$) (39).

17

18 The prevalence of ADA against adalimumab has high variability across different types of
19 autoimmune diseases in adults (25, 28, 29, 40-42) and children with JIA(36) (Table 1). The
20 timing of adalimumab ADA development is controversial: in some adult studies ADA
21 prevalence did not increase with treatment duration (43, 44), while in other studies there was a
22 significant increase, with ADA developing between 4.5 months and 12 months of treatment (9,
23 30, 40, 42, 45, 46). Similarly, studies in JIA showed both trends: a significant increase of ADA
24 with time (36) or no correlation with treatment duration (47), suggesting that ongoing
25 monitoring to establish their clinical relevance and impact on management is required.

26 Etanercept treatment was associated with a lower ADA rate than infliximab and adalimumab
27 (25) (Table 1), with the vast majority of adult studies reporting no detectable ADA (25, 27-29,
28 31, 40, 42, 46). This pinpoints that the chemical structure of the anti-TNF- α therapeutic agent
29 (fusion protein versus mAb) is likely to be a key factor in inducing drug immunogenicity. When
30 detected, ADA against etanercept were found to be non-neutralising in both adults and
31 paediatric studies (28, 36). ADA prevalence increased with treatment duration with a
32 corresponding decrease in etanercept drug levels over time in JIA (48, 49).

33

1 A highly sensitive ELISA test detected ADA against golimumab in 31.7% of patients with RA,
2 PsA and AS in comparison with standard ELISA which detected ADA only in 4.1% (50), while
3 their prevalence varied across adult studies (Table 1). The impact of ADA on serum golimumab
4 concentrations was consistent in JIA and RA studies, whereby higher ADA titers were
5 associated with lower drug concentrations (28, 51-53). This was generally shown at ADA titres
6 >1:1000 in JIA (51), and in adults, median peak titres ≥ 100 were associated with undetectable
7 or very low drug levels (59). Interestingly, in another study in PsA, which used a standard
8 assay, the golimumab dose (50mg vs. 100mg) did not appear to affect the ADA rates, which
9 remained low for the whole duration of the study through to week 52 (4.9%) (54).

10
11 There are fewer studies investigating the presence of ADA against certolizumab (55, 56),
12 although in both studies, ADA were associated with lower drug levels (Table 2). A more recent
13 study, however, reported that there was no significant correlation between ADA and
14 certolizumab drug levels ($r = -0.471$, $p = 0.122$). There is evidence that ADA were still detected
15 at higher certolizumab concentrations of >10mg/l (57). The majority of patients with ADA had
16 detectable titres from week 16 onwards and 65% remained ADA positive after one year of
17 follow up (57). There are no studies in paediatric populations.

18
19 When anti TNF- α agents have been studied comparatively in adults, there was evidence of
20 increased prevalence of ADA against infliximab compared to adalimumab (25.3% vs 14.1%
21 respectively), as well as between adalimumab and golimumab (14.1% vs 3.8%) (25). Similar
22 trend was found in a meta-analysis of biologic agents in JIA, where the pooled prevalence of
23 ADA against infliximab was 36.6% compared to 21.8% for ADA against adalimumab (36). As
24 mentioned above, the prevalence of ADA against golimumab seems to be higher in children
25 (46.8%) but based on limited evidence (51).

26
27 **Variable impact of ADA directed against anti TNF- α treatments on clinical efficacy:**
28 **loss of efficacy to adalimumab and infliximab was consistently found in children and**
29 **adults who developed ADA**

30 Various studies in RA, PsA, AS provided evidence for an association between the presence of
31 ADA against adalimumab and loss of clinical efficacy or diminished clinical response (23, 28,
32 29, 40), while other studies found no association (43, 44) (Table 1). The impact of ADA on the

1 trend of inflammatory markers is not clear; some studies found higher ESR and CRP in patients
2 who had detectable ADA (27, 29), whereas other studies found no such association (43). In
3 addition, the presence of both ADA and low adalimumab concentration at 3 months were
4 together significant predictors of poor response at 12 months (40, 42). However, the risk of
5 flares following various adalimumab tapering strategies in RA did not seem to be influenced
6 by the adalimumab serum levels or ADA prevalence (58).

7 A higher proportion of ADA positive JIA patients treated with adalimumab experienced loss
8 of response and more clinical relapses than those without ADA (28, 47). In JIA, it was noted
9 that transient ADA (defined as measurable ADA on up to two consecutive time points which
10 disappeared on subsequent measurements without having any impact on treatment efficacy or
11 toxicity) were not associated with diminished response to medication, whereas permanent
12 ADA did lower treatment response (45).

13

14 Most adult rheumatology studies found no detectable ADA against etanercept (27, 30). It has
15 been suggested that neither etanercept concentrations nor ADA positivity correlated with JIA
16 activity or remission states (48).

17

18 A meta-analysis of 9 studies of infliximab in adult autoimmune diseases found that the presence
19 of ADA decreased the odds of response by 58% (25). After 52 weeks of treatment with
20 infliximab, non-responder RA patients were significantly more likely to be ADA positive (34).

21

22 Adult RA studies found that ADA against golimumab were associated with a poorer clinical
23 response (28, 52). ADA positive RA patients (15.2% at 24 weeks) had a worse EULAR
24 response and higher DAS-28 compared to ADA negative patients (52). However, one study
25 which utilised a more sensitive method of ADA detection (drug-tolerant enzyme immunoassay,
26 DT-EIA) in adults, reported no effects of ADA to golimumab on clinical responses at 24 and
27 52 weeks, across RA, PsA and AS (50). This highlights the importance in sensitivities of assays
28 used. Studies in children with JIA found that ADA to golimumab did not appear to have impact
29 on clinical responses (59) (51). Brunner et al., reported that none of the 8 JIA patients found
30 with high ADA titres >1:1000, experienced flares (51).

31

32 ADA against certolizumab appeared to have an impact on RA clinical response at 3 months,
33 where the majority of ADA positive patients were non-responders (56), but there was no

1 independent correlation with the 12 month EULAR response (55), suggesting that there was a
2 time-dependent relationship. There are no paediatric studies.

3

4 A meta-analysis performed on 12 observational prospective cohort studies in adults evaluated
5 that the development of ADA reduced the anti-TNF response rate (RR) by 68% (RR = 0.32,
6 95% CI 0.22, 0.48)(60), while in children with JIA, a qualitative analysis found that antibodies
7 to infliximab and adalimumab were associated with treatment failure (36).

8

9 **Additional methotrexate treatment decreased the rate of ADA formation against anti**
10 **TNF- α treatments**

11

12 Generally, for both adults and children, concomitant DMARD therapy was beneficial and
13 resulted in a decrease in ADA positivity, but the impact of DMARDs on ADA formation was
14 not always analysed to enable reliable conclusions (9, 47) (Table 1). Most studies looked at
15 concomitant methotrexate (MTX) therapy but azathioprine, leflunomide and mycophenolate
16 have also been shown to be associated with lower ADA prevalence, suggesting that all
17 DMARDs may be associated with benefits against drug-induced immunogenicity (23, 28, 42)
18 (31). Unfortunately, none of the studies evaluated comparatively the impact of individual
19 DMARDs on immunogenicity in inflammatory arthritis because of small numbers of patients
20 on DMARDs other than MTX, and because some patients were treated with more than one
21 conventional DMARD. Concomitant use of MTX was associated with lower rates of ADA
22 against infliximab in RA (28, 31, 32, 40, 61). Moreover, RA patients treated with infliximab
23 were less likely to develop ADA if they received high biologic doses/induction therapy, or if
24 they received continuous versus intermittent therapy (28, 30, 32, 61, 62). A RCT of infliximab
25 plus MTX for the treatment of JIA, found that more patients achieved clinical response in the
26 ADA negative group (79% vs. 67%) (37).

27 Similar evidence has been found in children, with studies suggesting a protective effect with
28 the addition of MTX (36, 45, 59). Interestingly, DMARD use in children was found to be
29 significantly lower in those who developed permanent ADA to adalimumab (45). It has also
30 been suggested that MTX reduces immunogenicity against adalimumab in a dose dependent
31 manner (30, 40), as patients who did not develop ADA were on a higher MTX dose (46).
32 However, a paediatric study found that there was no difference in ADA rates in JIA patients
33 with longer exposure to MTX (47).

1 In adults, concomitant use of MTX was associated with lower incidence of ADA to golimumab
2 (28, 50, 63). A study found that the mean trough golimumab level at 24 weeks was comparable
3 in ADA positive vs. negative patients, with or without concomitant MTX (63).

4

5 **ADA against infliximab and adalimumab have been associated with side-effects to**
6 **therapy**

7

8 In both adults and children, there was no clear consensus on whether ADA have an impact on
9 safety (Table 1). As expected, most reports included a small number of cases experiencing
10 side-effects. Adverse events more frequently mentioned included injection site or infusion
11 reactions, serum sickness and thromboembolic events. Some studies suggested that adverse
12 events occurred more frequently in patients with ADA to adalimumab (28, 29, 62) with others
13 showing no significant differences (27, 44). In paediatric studies, despite limited information
14 available, no association between the presence of ADA and adverse events was reported (36).
15 There was a suggestion of a possible increase in minor upper respiratory tract infections in
16 children with detectable ADA, however, this conclusion was limited by the small sample size
17 (45).

18 ADA against infliximab have been reported to confer a higher likelihood of adverse drug
19 reactions (25, 28, 30, 32, 35, 40, 62). In a RA study (35), ADA positive patients had an
20 increased risk of adverse drug reactions compared with ADA negative patients over 52 weeks
21 [21 (18%) vs. 7 (7%), $P < 0.018$] (40). Similarly, JIA infusion reactions to infliximab were
22 more commonly seen in ADA positive patients (58% vs 19%) (37). A retrospective chart
23 review of children with JIA and paediatric inflammatory ocular diseases found that patients
24 with ADA had a 15-fold increased risk of infusion reactions to infliximab compared to patients
25 without ADA (39). This study also found that ADA positive children were significantly
26 younger (mean age 7.01 vs. 9.88 years, $p = 0.003$).

27

28 Limited data were available regarding the impact of immunogenicity against etanercept on
29 safety. Studies across age did not report an association between ADA positivity and adverse
30 events (36, 59). In JIA studies, the proportion of patients with ADA did not differ between
31 responders and non-responders to etanercept (48).

32

1 Studies in both paediatric and adult populations did not report an association between ADA
2 and adverse effects to golimumab (51, 52, 59). Similarly, multiple adult studies reported no
3 association between the presence of ADA against certolizumab and adverse effects (55-57); in
4 addition, RA patients who experienced adverse effects did not have ADA (55, 56).

5
6
7 **Immunogenicity to anti TNF- α biosimilars is similar to or lower than that of their**
8 **originators**

9
10 Biosimilars are new biological products which are highly similar to their biological reference
11 drug and have comparable clinical efficacy. At present, the use of biosimilars in JIA is limited,
12 thus the majority of evidence related to their immunogenicity is available from adult studies.
13 Multiple studies have shown similar clinical efficacy and immunogenicity profiles when
14 comparing biosimilars with their reference products (28, 64-72). For example, ADA positive
15 CT-P13 (an infliximab biosimilar) patients showed less clinical improvement (28). ADA
16 against infliximab and adalimumab biosimilars were associated with lower drug concentrations
17 (69)(75). The PLANETRA study found that peak serum CT-P13 concentrations were reduced
18 in the ADA positive group ($C_{max} = 85.1\mu\text{g/ml}$) compared to the ADA negative subset ($C_{max} =$
19 $96.7\mu\text{g/ml}$) (69). One meta-analysis reported on the pooled response rates (RR) of ADA
20 against anti TNF- α biosimilars compared to their reference product (66). There were no
21 significant differences in ADA formation rates between the infliximab and adalimumab
22 biosimilars and their reference drugs at 24-30 weeks. The etanercept biosimilars showed
23 significantly lower rates of ADA formation compared to the reference product, with a pooled
24 $RR = 0.05$ at 24-30 weeks (66). A study of etanercept biosimilar GP2015 did not detect any
25 neutralising ADA, and all ADA responses were transient (absent by week 24) (72).

26
27
28 ***Clinical relevance of ADA against other biologic agents in adult and paediatric***
29 ***inflammatory arthritis studies***

30
31 **ADA against abatacept are mainly non-neutralising and do not have significant impact**
32 **on clinical efficacy unless treatment is temporarily discontinued**

1 The prevalence of ADA to fusion proteins, such as abatacept (which comprises a Fc region of
2 IgG1 fused to the extracellular domain of CTLA-4) is generally acknowledged to be lower than
3 to therapeutic mAbs. The prevalence of ADA to abatacept ranged from 1-20% in adult studies
4 (28, 30, 41, 73), and from 8.7-23.3% in paediatric studies (36) (Table 2). Younger children
5 with JIA (2-5 years) had a higher prevalence of ADA than older children (6-17 years) (74).
6 One JIA study compared the prevalence of abatacept specific ADA with anti- CTLA-4 specific
7 antibodies and found the latter to be much higher (1.2% vs. 20.7%) (75). In terms of timing of
8 the development of ADA in children, one study found that ADA concentration increased with
9 a longer time of exposure to abatacept (76), whereas another found no increase with continued
10 exposure (77).

11 Similar to etanercept, abatacept generated ADA which bind to the Fc fragment (hinge region)
12 and have no neutralising activity (28). Non-neutralising ADA decreased the circulating levels
13 of abatacept by enhancing drug clearance in adults (30, 41). In children, ADA were also found
14 to be non-neutralising but were not found to be associated with low abatacept concentrations
15 (75, 76).

16 No loss of efficacy due to ADA against abatacept was found in JIA studies (36, 75-77), while
17 in contrast, in adults with RA, intermittent treatment discontinuation led to higher incidence of
18 immunogenicity and loss of clinical response (73). It was observed that adult patients who
19 discontinued the treatment temporarily had a higher ADA rates than those on continuous
20 treatment (7.4% vs 2.6% respectively) (30). Similarly, ADA were more frequent in children
21 with JIA who interrupted treatment and had abatacept concentration below therapeutic levels,
22 suggesting that higher treatment doses may be beneficial against immunogenicity (75).

23 Some adult studies suggested that intravenous therapy was associated with less
24 immunogenicity than subcutaneous administration (28),(78), while other studies found no
25 difference (30). In JIA, no difference was found between the two routes of administration (36).

26

27 In RA, concomitant MTX therapy did not significantly affect immunogenicity (73). In
28 paediatric studies the impact of MTX has not been studied (36). Reassuringly, ADA against
29 abatacept were not associated with increased risk for injection site reactions, hypersensitivity
30 or any other safety concerns (36, 73, 75, 76), even when patients have been followed up to 7
31 years (77).

32

1 **ADA against B cell targeted therapies are dose-dependent and have impact on clinical**
2 **efficacy and risk of adverse reactions**

3
4 Rituximab is a chimeric mAb against CD20. There have been no paediatric studies
5 investigating the relevance of ADA against rituximab. However, ADA against rituximab have
6 been reported in 0-21% of adult RA patients (28). Additionally, ADA have been found to be
7 associated with a reduced treatment response and higher rates of treatment serious adverse
8 events (28, 79). Lower serum rituximab concentrations have been reported in ADA positive
9 patients compared to ADA negative patients in RA (80). Moreover, the use of higher rituximab
10 doses and induction therapy have been associated with a decreased incidence of ADAs in RA
11 (28).

12
13 A meta-analysis reported that the pooled RR of ADA formation for rituximab biosimilars was
14 0.86 at week 24-28 (67). Of note, the pooled RR of neutralising ADA formation at the same
15 time point was 1.16. Neutralising ADA were also of a very low incidence at week 72 in the
16 rituximab biosimilar CT-P10 (68). Multiple studies have demonstrated a similar side effect
17 profile for biosimilars, as higher rates of infusion-related reactions were present in ADA-
18 positive patients compared to ADA-negative patients (28, 64, 65, 70, 71) (Table 2).

19
20 **Neutralising ADA against tocilizumab have no clear impact on clinical efficacy and**
21 **potential on side-effects in adults, while there is a trend for clinical impact in children**

22
23 Tocilizumab is a humanized mAb against the interleukin-6 receptor (IL-6R). Several studies
24 have reported low ADA rates in RA patients (28) (81, 82). ADA positivity has been recorded
25 in 1.5% and 1.2% of RA patients receiving intravenous and subcutaneous tocilizumab
26 respectively, with a high proportion of these being neutralising ADA (83) (Table 2). The rate
27 of ADA formation has not been seen to significantly differ in tocilizumab monotherapy versus
28 combination therapy with conventional synthetic DMARDs (83). No correlation has been
29 found between ADA rates and adverse events or a reduced treatment efficacy in adults (41,
30 83). Similarly, low levels of ADA to tocilizumab have been reported in JIA patients, with a
31 pooled prevalence of 2.3% across four studies (36). However, neutralising antibodies against
32 tocilizumab in JIA have indeed been shown to correlate with treatment failure, as well as with
33 infusion and hypersensitivity reactions (36, 84). Yokota et al. (84) found that out of five JIA

1 patients treated with tocilizumab who developed ADA, four (80%) withdrew from the study
2 due to infusion reactions.

3
4 **ADA to sarilumab seem to have limited impact on clinical efficacy and no impact on**
5 **adverse events**

6
7 Sarilumab is human recombinant mAb that blocks both the soluble and membrane-bound IL-
8 6 receptor, similarly to tocilizumab, but with a higher affinity. Currently there are no studies
9 of immunogenicity in paediatric populations. The presence of ADA did not appear to affect
10 clinical efficacy in various trials (85-87). The MONARCH trial demonstrated that only 2.7%
11 of RA patients had persistent ADA, however, no neutralising ADA were detected (85). It has
12 been suggested that ADA against sarilumab are in majority of cases transient (88). Xu et al.
13 described a trend towards higher apparent linear clearance of sarilumab when ADA were
14 present (89). In addition, patients with persistent ADA had a lower mean drug levels compared
15 to ADA negative patients. At a dose of 150mg, treatment-emergent ADA incidence was 24.6%
16 compared to 18.2% at a higher dose of 200mg. Of those who had persistent ADA, the incidence
17 of neutralising ADA was also higher in the group receiving 150mg sarilumab compared to
18 200mg (10.8% and 3.0% respectively) (86). Multiple studies have shown that ADA positivity
19 was not associated with a higher incidence of adverse effects (85) (86, 87). Hypersensitivity
20 reactions occurring during treatment were reported in 8.0% of ADA-negative patients and in
21 3.1% of ADA-positive patients (87).

22
23 **Neutralising ADA against IL12/23 blockade have low prevalence but possible impact on**
24 **clinical efficacy in inflammatory arthritis**

25
26 Ustekinumab is a human immunoglobulin G1 κ monoclonal antibody against common sub-unit
27 p40 of IL-12 and IL-23. The prevalence of ADA was 8-11% in psoriatic arthritis adult patients
28 treated with ustekinumab (28). Moreover, a study evaluating the efficacy of subcutaneous
29 ustekinumab in the treatment of RA reported that 7/123 (5.7%) of patients had ADA, while
30 4/123 (3.3%) had neutralising ADA (90). In this study, serum concentrations of ustekinumab
31 were generally lower in ADA positive patients (90) (Table 2). There is evidence that
32 neutralising ADA against ustekinumab were associated with lower drug levels and loss of
33 clinical efficacy in psoriasis and Crohn's disease (91, 92), suggesting overall that they may

1 have similar impact in inflammatory arthritis. The relevance of ustekinumab immunogenicity
2 is yet to be studied in children.

3
4

5 **Very low prevalence of ADA against IL-17 blockade has been reported and no impact**
6 **on side-effects or clinical efficacy**

7

8 Secukinumab is a mAb targeting IL17A. The treatment is not licensed for children. In a recent
9 systematic review, the prevalence of ADA against secukinumab was 0-1% (28). A study
10 evaluated the prevalence of ADA at 52 weeks in patients with psoriasis, PsA and AS treated
11 with secukinumab and found it to be <1%; ADA were not associated with loss of efficacy,
12 changes in drug levels or adverse events (93).

13 Ixekizumab is a humanized mAb which targets IL17A used for the treatment of plaque
14 psoriasis, PsA and AS. The prevalence of ADA was 5.3% (94) and 9% (95) in adult patients
15 with psoriasis and PsA, and they occurred within the first 12 weeks of treatment (95). ADA
16 were found to be non-neutralising and did not correlate with the rate of adverse reactions (Table
17 2). Patients with psoriasis or PsA who developed ADA against ixekizumab had low and
18 constant titres, which did not significantly impact clinical response. No data in children are
19 available.

20 **ADA against IL-1 blockade do not have significant impact on clinical efficacy or side-**
21 **effects**

22

23 Anakinra is a recombinant a human IL-1 recombinant receptor antagonist initially trialled in
24 RA, where it has been associated with a prevalence of ADA ranging from 50.1 to 70.9% (96,
25 97). Similar to other recombinant proteins, only a small proportion of ADA were neutralising
26 (25/1240, 1.9%) (96) (Table 2). Of these 25 RA patients, 13 (52%) reported disease
27 progression; however, no relationships were found between neutralising antibody status and
28 the occurrence of severe allergic reactions, malignancies, opportunistic infections, or serious
29 infections (96). One study assessing the efficacy of anakinra in patients with JIA found that
30 the prevalence of ADA increased from 75% at 12 weeks to 82% at 12 months (98). At 12
31 weeks, all 4/64 (6%) of patients who had neutralising antibodies to anakinra were non-
32 responders to treatment (98). However, non-neutralising antibodies to anakinra were not

1 associated with a reduced response to treatment (98). There have been no studies analysing the
2 association between ADA to anakinra and adverse events in JIA.

3

4 Canakinumab is a fully human mAb against anti-IL1- β used in systemic-onset JIA (soJIA).
5 Studies in children with systemic JIA found a prevalence of ADA against canakinumab of
6 3.1% (6/196) (99), and 8% (100), and ADA had no neutralising capacity and did not affect the
7 drug levels or the rate of side-effects.

8

9 Rilonacept is a fully human dimeric fusion protein that acts as a soluble decoy receptor which
10 blocks IL-1 β . An RCT in soJIA did not find an association between ADA positivity and clinical
11 response (101). This trial found that 54.2% (13/24) of patients developed ADA during the 23-
12 month period of open label treatment (following a 4-week double blind treatment phase). There
13 was no correlation between ADA positivity and plasma levels of rilonacept (101). Although
14 the sample size was small, this study noted that the patients who developed ≥ 3 injection site
15 reactions were all ADA positive, thus suggesting that there is an association between ADA and
16 adverse effects.

17

18 **Conclusion:**

19

20 Immunogenicity to biologic treatment has been investigated in various types of inflammatory
21 arthritis in children and adults. The overall impression is that immunogenicity to biologics used
22 in rheumatology was not particularly confounded by clinical indication or significantly affected
23 by patients' age (Table 3). However, a direct comparison between the studies evaluated by this
24 report is not possible, because of the high study heterogeneity, low number of studies
25 investigating less commonly used biologic treatments and high variability between the methods
26 of ADA detection and time-points of ADA measurements, study design and concomitant MTX
27 therapy.

28

29 As there are some differences between the biologic agents approved for use in paediatric versus
30 adult rheumatic diseases, in some cases there were no data available to enable comparisons
31 between the two populations (e.g. certolizumab, sarilumab, secukinumab, ustekinumab and
32 ixekimumab have no studies in children, while rilonacept and canakinumab are not commonly

1 used in adults). The discrepancy found between the rate of ADA against golimumab is not easy
2 to interpret, because they have been investigated only in one study in JIA.

3 This literature review provided evidence for variable prevalence of ADA depending on the
4 study methodology, sample size, time-points for sample evaluation, concomitant DMARD
5 therapy as well as laboratory assays used for ADA detection. Overall, the highest ADA
6 prevalence was found in patients treated with mAbs against TNF- α and recombinant human
7 IL-1 receptor antagonist (anakinra), although the impact of ADA on clinical efficacy was
8 clearly influenced by their neutralising properties and impact on drug levels. In contrast to
9 immunogenicity to IL-1 blockade, which had minimal or no impact on clinical efficacy as the
10 proportion of neutralising ADA was very low, ADA against adalimumab, infliximab,
11 certolizumab, and to a certain extent golimumab had a significant impact on clinical efficacy.
12 As a consequence, the choice of biologic therapeutic agent in a certain patient influences their
13 immunogenicity monitoring strategy.

14

15 All mAbs against TNF- α (and their biosimilars) were associated with higher prevalence of
16 ADA than etanercept (a fusion protein) and this is probably explained by the structure of the
17 biologic agent as well as frequency of administration, which in the case of etanercept ensures
18 a more constant serum drug levels. It is recognized that anti-idiotypic ADA against therapeutic
19 mAbs usually target the drug binding site as this does not belong to the patient immunoglobulin
20 repertoire, therefore these ADA have neutralising properties with impact of drug efficacy and
21 they are clinically relevant (62). The detection of neutralising ADA in certain patients should
22 be monitored and correlated with clinical response and drug levels to guide further therapeutic
23 decisions (102). Neutralising ADA have been found in patients treated with adalimumab,
24 infliximab, certolizumab pegol, and golimumab, as well as tocilizumab, ustekinumab and
25 secukinumab.

26

27 By contrast, in the case of fusion proteins which comprise a naturally occurring receptor fused
28 with the constant region of human Ig, the immunogenicity process is primarily triggered by the
29 recognition of the fusion part of the molecule with no direct impact on the drug binding site.
30 Overall these therapeutic agents were associated with less immunogenicity, although
31 neutralising ADA against fusion proteins have also been described with both etanercept and
32 abatacept (103, 104), suggesting that their monitoring could be relevant in selected categories
33 of patients, especially if the treatment has been discontinued temporarily.

34

1 Despite the potential side-effects associated with the presence of ADA overall, irrespective of
2 their neutralising properties, detection of ADA does not preclude loss of clinical response as
3 long as it does not reduce the serum concentration of the biologic agent below the therapeutic
4 threshold (62), therefore monitoring of ADA without drug levels has no clinical relevance.

5

6 High ADA concentration correlated with lower drug levels and impact on clinical efficacy
7 when patients of all ages were treated with adalimumab, infliximab, golimumab, certolizumab,
8 rituximab, abatacept, anakinra, canakinumab, and possibly ustekinumab, while the presence of
9 ADA had less impact on clinical efficacy in adult patients treated with IL-6 and IL-17 blockage
10 and children treated with rilonacept (IL-1 β decoy receptor). Patients with higher ADA titers
11 and lower or not/detectable drug levels are probably at risk of losing clinical efficacy and need
12 to be monitored more closely.

13

14 It is clinically important to take into consideration the fact that not all detectable neutralising
15 ADA had impact on clinical outcomes (e.g. tocilizumab ADA lowered treatment response in
16 children with JIA but less in adults with RA). Neutralising ADA were more commonly found
17 in patients treated with mAbs compared to fusion proteins; however, not all ADA against mAbs
18 had neutralising properties or impact on clinical efficacy (e.g. ADA against ixekizumab were
19 predominantly non-neutralising and did not influence clinical response).

20

21 The timing of developing ADA varied according to the type of biologic treatment and patients'
22 age. Patients developed ADA against adalimumab earlier in their disease course, while ADA
23 in children with JIA treated with abatacept increased with longer time exposure to the drug.

24 Although data from paediatric studies are scarce overall, studies found that younger age in
25 children with JIA was associated with a higher prevalence of ADA as well as side-effects to
26 certain biologics, suggesting that caution in monitoring younger patients is advisable.

27

28 There is good evidence that higher doses of rituximab and infliximab, as well as more regular
29 administration (as in the case of etanercept) were associated with lower ADA prevalence,
30 suggesting that medication discontinuation and tapering biologic treatment doses could have
31 impact on clinical efficacy. Monitoring patients' compliance and taking into consideration their
32 dosing regimen, route and frequency of biologic medication administration are important
33 aspects of immunogenicity risk assessment. Increasing treatment dose as well as switching to

1 IV formulations can lower the ADA and restore treatment response, therefore these are useful
2 therapeutic strategies to address the clinical impact of drug-induced immunogenicity.

3

4 In addition, the large variability of ADA levels against biologic agents detected in various
5 adult and pediatric studies of inflammatory arthritis is very likely influenced by the sensitivity
6 of the assay used, concomitant MTX dose, time point of sample collection, as well as patients'
7 characteristics (genetic background, smoking, age). The overall impact of ADA on drug
8 efficacy, as well as therapeutic drug monitoring are particularly relevant in guiding future
9 therapeutic strategies of tapering biologic treatments in inflammatory arthritis patients (102,
10 105), although further research related to their impact on clinical decision making is required
11 (16, 58).

12

13 Based on data available in the literature, concomitant treatment with MTX to address the risk
14 of immunogenicity is recommended in patients treated with abatacept, infliximab, golimumab,
15 while in the case of treatment with etanercept, abatacept and tocilizumab the impact of
16 additional MTX is not significant.

17

18 We propose a potential strategy for drug immunogenicity monitoring for improved clinical
19 benefit (Figure 1). The main clinical instances when ADA and drug levels should be monitored
20 is loss of clinical efficacy, monotherapy with biologic agents recommended to be prescribed in
21 addition to MTX, clinical reasons for frequent dose intermittent discontinuation, in patients
22 who tapered biologics (especially administered subcutaneously), patients who develop
23 infusion/injection reactions and other side-effects to therapy. Further research especially
24 focused on patient individual risk to develop immunogenicity to biologics is required to enable
25 personalized therapy selection.

26

27

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Author et al., year [ref]	Country Type of study (including meta-analyses) Number of patients treated with a certain biologic	Type of inflammatory arthritis Age range or mean age (years)	Disease duration Range or mean± SD (years)	Prevalence of ADA Impact of additional DMARD therapy on ADA prevalence	Impact on clinical efficacy	Impact on side-effects to biologic therapy
Adalimumab and biosimilars						
Strand et al.,2017 [28]	Systematic review RA N= 1282 PsA N= 59 JIA N=23 AS = 204	RA (35-64) PsA (43-55) JIA (3-14.2) AS (30-48)	RA: 1-34 PsA: 5-21 JIA: 1-5 AS: 4-15	RA 0-51%; PsA 0-54% JIA 6-33%; AS 8-39% Concomitant use of MTX, AZA, leflunomide or MMF was associated with lower rates of ADA in RA, JIA, AS	ADA was associated with less improvement of disease activity for RA, PsA and AS. A higher proportion of ADA+ve JIA patients experienced loss of response than ADA-patients (no P value reported).	Adverse events occurred more frequently in ADA+ve patients compared to ADA-ve (27% vs 15%, no P value reported)
Doeleman et al.,2019 [39]	Systematic review and meta-analysis N= 355	JIA 10.5	3.45	Pooled prevalence of 21.5% (95% CI = 14.1 – 29.8) Addition of MTX reduced the risk of ADA development by 67% (RR 0.33)	Increased median disease activity score in patients with ADA was found (no P value reported)	No association with adverse events generally was found, but in patients with JIA-associated uveitis, ADA were associated with a significantly higher severity of uveitis (no P value reported).
Marino et al., 2018 [47]	Italy Prospective observational study N=27	JIA Age at inclusion 9.5±3.32 ADA+ve 11.15 ± 3.11	4.79± 3.04	Overall prevalence 37% 31% vs. 45% in MTX+ve vs. MTX-ve groups. No impact of MTX treatment duration on ADA development was found -22.9 months	ADA+ve patients experienced more relapses, P<0.017. 30% of ADA+ve patients were in clinical remission,	No infusion reactions or side effects were found

		ADA-ve 8.52 ± 3.12		(MTX+ve group) vs. 17.8 months (MTX-ve group)	compared to 41.2% of ADA- patients, P=0.56	
Maid et al.,2018 [29]	Argentina Cross-sectional study N=52	RA 56.5 (13.3)	10.8± 8.5	36.5%. 36% of MTX+ve patients and 38% of MTX-ve patients tested positive for ADA	ADA-ve patients had a tendency towards better clinical outcomes than those who were ADA+ve – 39.4% of ADA-ve patients achieved an HAQ-DI score <0.5, compared to only 31.6% of ADA+ve patients (comparative statistics were not performed)	Injection site reactions were reported by 6.3% in the ADA-ve group and 4.3% in the ADA+ve group (no p-value reported) (combined data for adalimumab, infliximab and etanercept)
Balsa et al., 2018 [31]	Spain Cross-sectional, observational study N=217	RA and SpA RA = 56.3 (12.1) SpA = 47.9 (11.5)	RA = 13.9 ± 8.7 SpA = 12.5 ± 10.2	RA: 25.5%; SpA: 32.7% No significant difference between the two patient groups (p=0.221) Lower proportion of patients receiving concomitant DMARDs (24.1% vs 36.9% were ADA+ve, P=0.037).	82.5% ADA+ve patients had no detectable drug levels in the serum. Only one ADA+ve patient reported drug concentrations within the normal range. No p-value reported.	Data not available
Quistrebert et al., 2019 [9]	European retrospective multi-cohort analysis N=240	RA 50.3	2.18	19.2% 96.6% of patients were MTX+ve, but study was not powered to analyse the effects	ADA positivity was significantly associated with a lower probability of a good clinical response based on 278 clinical observations from 215 patients (hazard ratio = 0.58, 95% CI 0.39–0.86)	Data not available
Verstegen et al., 2020 [62]	Systematic review N= 103	JIA 10.6	Data not available	6.7%-37% Concomitant treatment with MTX showed a protective effect against ADA development for	ADA to adalimumab were associated to impaired clinical outcome (no comparative statistics performed)	Data not available

				patients treated with adalimumab and infliximab		
Skrabl-Baumgartner et al., 2019 [45]	Austria Prospective observational study N=20	JIA 9.9± 4.2	JIA data not available Duration of JIA-associated uveitis 3.5+/-3.5	45% (including permanent and transient ADA) Concomitant use of DMARDs significantly lower in group with permanent ADA+ve (2/7) vs ADA-ve (10/11) – p<0.05	7/8 who had a loss of response had permanent ADA. Transient ADA were not associated with a diminished response (no comparative statistics performed)	No severe adverse reactions were found.
Moots et al., 2017 [27]	Multinational non-interventional study N=199	RA 54.3± 12.95	Symptom duration 9.3 ± 8.43	RA 31.2%	Significant differences between patients with and without detectable ADA were observed in ESR (p=0.008) and CRP (p=0.0011). When data for all three TNF inhibitors were pooled, a greater proportion of patients without detectable ADA (226/484; 46.7%) than those with detectable ADA (29/94; 30.9%) were in remission (p=0.0046).	No differences in safety outcomes were reported
Infliximab and biosimilars						
Strand et al., 2017 [28]	Systematic review RA N=1412 PsA N= 173 JIA N = not available AS N = 163	RA (35-64) PsA (43-55) JIA (3-14.2) AS (30-48)	RA: 1-34 PsA: 5-21 JIA: 1-5 AS: 4-15	RA 8-62%; PsA 15-33%, JIA 26-42%; AS 6.1-6.9%; Concomitant use of MTX, AZA, leflunomide or MMF is associated with lower rates of ADA in RA	ADA+ve patients showed less improvement in disease activity and were less likely to achieve clinical responses (RA, PsA, AS) - (no comparative statistics performed)	Increased risk of treatment discontinuation due to adverse events and higher rates of infusion reactions were reported in ADA+ve patients (no comparative statistics performed)

Maid et al.,2018 [29]	Argentina Cross-sectional study N=13	RA 55.5 (10.6)	13.1±8.5	30.8% 22.2% of MTX+ve and 50% of MTX-ve patients had ADA	ADA-ve patients had a tendency towards better clinical outcomes than those who were ADA+ve – no comparative statistics were performed due to low numbers.	Injection site reactions were reported by 6.3% in the ADA-ve and 4.3% in the ADA+ve group (no p-value reported.)(combined data for adalimumab, infliximab and etanercept)
Balsa et al., 2018 [31]	Spain Cross-sectional, observational study N= 188	RA and SpA RA = 56.3 (12.1) SpA = 47.9 (11.5)	RA = 13.9 ± 8.7 SpA = 12.5 ± 10.2	RA: 21.1%; SpA: 31.3% No significant difference between the two patient groups (p=0.114) Concomitant use of DMARDs associated with lower ADA – ADA-ve 29/130 (22.3%) vs 22/58 ADA+ve (37.9%); P = 0.021)	78.4% ADA positive patients had no detectable drug in the serum. Only one ADA+ve patient reported drug concentrations within the normal range. No p-value reported.	Data not available
Quistrebert et al., 2019 [9]	European retrospective multi-cohort analysis N=126	RA 50.6	2.65	RA 29.4% ADA were detected more frequently in infliximab-treated patients (29.4%) than in adalimumab-treated patients (19.2%).	ADA positivity was significantly associated with a lower probability of a good clinical response based on 149 clinical observations from 125 patients(hazard ratio = 0.61, 95% CI 0.32–0.76)	Data not available
Ruperto et al., 2007 [36]	Multicentre RCT N=122	JIA 11.2	3.9	25.5%	Data not available	Infusion reactions were observed in 58% of ADA+ve patients compared to 19% of ADA-patients. Serious infusion reactions additionally occurred in 20% of ADA+ve patients,

						compared to 0% of ADA-patients. No comparative statistics performed
Ruperto et al., 2010 [37]	Multicentre open-label extension study N= 78	JIA Data not available	Data not available	37% (+32% inconclusive)	Data not available	32% patients had ≥1 infusion-related reaction, with a higher occurrence amongst patients who were ADA+ve (15/26 [58%] ADA+ve patients had infusion-related reactions). No comparative statistics performed
Moots et al., 2017 [27]	Multicentre noninterventional study N=196	RA 60.7±13.01	Symptom duration 10.0±10.11	RA 17.4%	95/184 (51.6%) were in low disease activity, of which 14/32 (43.8%) had detectable ADA and 81/152 (53.3%) had no detectable ADA (P = 0.3387). Significant differences between patients with and without detectable ADA were observed in ESR (p<0.0001) and CRP (p=0.0001).	No significant correlation between adverse events and ADA was found.
Etanercept and biosimilars						
Strand et al.,2017 [28]	Systematic review RA N=589 PsA, JIA, AS N = not available	RA (35-64) PsA (43-55) JIA (3-14.2) AS (30-48)	RA: 1-34 PsA: 5-21 JIA: 1-5 AS: 4-15	RA 0-13%; PsA 0% JIA 0-6%; AS 0%;	Data not available	Data not available

Balsa et al., 2018 [31]	Spain Cross-sectional, observational study N= 165	RA and SpA RA = 56.3 (12.1) SpA = 47.9 (11.5)	RA = 13.9 ± 8.7 SpA = 12.5 ± 10.2	RA: 0%; SpA: 0%	Data not available	Data not available
Doeleman et al., 2019 [39]	Systematic review and meta-analysis N= 268	JIA 11.8	4.7	Pooled prevalence 8.5% (95% CI = 0.5 – 23.2)	No reported association between treatment failure and the presence of non-neutralizing ADA	No association between adverse events and ADA was observed
Maid et al., 2018 [29]	Argentina Cross-sectional study N=54	RA 54.5 (13.6)	12.5±10.1	0%	Data not available	Data not available
Bader-Meunier et al., 2019 [48]	France Prospective multi-centre study N=126	JIA 10.5 (2-17)	4.62 (0.16-16.3)	15.7% at baseline 33% after 366 (302-712) days of treatment	ADA levels not significantly different between responders and non-responders (7.22±3.60 vs. 6.47±3.98ng/ml), No significant difference with concomitant MTX. p-values < 0.05 were considered significant.	No severe adverse events occurred.
Moots et al., 2017 [27]	Multicentre non-interventional study N=200	RA 56.5±13.37	Symptom duration 0.8±10.67	0%	No patients developed ADA on ETN).	Data not available
Constantin et al., 2016 [49]	Multicentre prospective open-label study	JIA 8.6± 4.6 ERA 14.5± 1.6 JPsA 14.5±2.0	JIA 31.6±31.7 months ERA	JIA - 18.3%, ERA- 23.7%, JPsA 20.5%, combined - 20.7%	No significant changes in effectiveness in patients who were ADA+ve was found	No safety concerns in patients who were ADA+ve were reported

	N=127		23.0±19.8 months JPsA 21.8±20.2 months	None of the ADA+ve patients had neutralising antibodies		
Golimumab						
Strand et al.,[28]	Systematic review RA N=1249 PsA, JIA and AS N = not available	RA (35-64) PsA (43-55) JIA (3-14.2) AS (30-48)	RA: 1-34 PsA: 5-21 JIA: 1-5 AS: 4-15	RA: 2-10%; PsA: 6%, AS: 0-6.4% Concomitant use of MTX, AZA, leflunomide or MMF was associated with lower rates of ADA in RA, PsA and AS	ADA+ve RA patients showed less improvement in disease activity and were less likely to achieve clinical responses (no comparative statistics performed)	Data not available
Brunner et al, 2018 [51]	Multicentre withdrawal RCT N=154	JIA 11.1+/-4.5	Disease duration not available	46.8% (72/154)	ADA did not appear to have a substantial impact on clinical efficacy	ADA were not associated with injection site reactions, disease flares or adverse events
Leu et al., 2019 [50]	Samples from 3 RCTs	RA PsA AS	Data not available	RA: 24.9% PsA: 39.9% AS: 30.3%	No effect of ADA on clinical response was found	Injection-site reactions were not affected by ADA
Kneepkens et al, 2014 [53]	The Netherlands Prospective observational cohort study N=37	RA	Data not available	8.1%	3 patients out of 37 (8.1%) were ADA+ve at 52 weeks and all 3 discontinued golimumab prematurely due to inefficacy	Data not available
Certolizumab						
Strand et al.,2017 [28]	Systematic review RA N= 358 PsA, JIA and AS N = not available	RA (35-64) PsA (43-55) JIA (3-14.2) AS (30-48)	RA: 1-34 PsA: 5-21 JIA: 1-5 AS: 4-15	RA 2.8-37% Concomitant use of MTX, AZA, leflunomide or MMF was associated with lower rates of ADA	Data not available	Data not available

Gehin et al.,2019 [56]	Norway Longitudinal observational study N=116	RA, AS, PsA and other inflammatory joint disease 42	2.6 0.6-14.1	Prevalence 6.1% (19/310 patients: 6 AS, 5 RA, 4 PsA and 4 other IJD) Among RA patients, 80% of ADA+ve patients had concomitant synthetic DMARDs (mostly MTX) vs. 73% of ADA- patients.	9% ADA+ve patients were responders at 3 months vs. 55% of ADA- patients No p- value reported	Data not available 8 patients experienced one or more injection-site reactions, all of which were ADA- at 3 months.
Jani et al., 2017 [55]	The Netherlands Prospective observation cohort study N=115	RA 58.0 ADA+ve 57.3 ADA-ve 58.5	7.0 3.3-14.4 ADA+ve 8.3 ADA-ve 6.0	37%	No correlation between ADA+ve and EULAR response was found (p = 0.18)	Data not available

Table 1 - Impact of ADA on disease outcomes in children and adults with inflammatory arthritis treated with anti TNF- α agents.

Legend: ADA- antidrug antibodies; AS – ankylosing spondylitis; AZT – azathioprine; ERA - enthesitis-related arthritis; EULAR- European League Against Rheumatism; JIA- juvenile idiopathic arthritis, JPsA – juvenile psoriatic arthritis; MMF- mycophenolate mofetil; MTX- methotrexate; N – number of patients treated with a certain biologic included in the study/systematic review; RA- rheumatoid arthritis, RCT – randomised control trial; PsA- psoriatic arthritis; +ve – positive; -ve - negative

<i>Author et al., year [ref]</i>	<i>Country Type of study</i>	<i>Type of inflammatory arthritis N (F:M) Age (mean+/-SD)</i>	<i>Disease duration</i>	<i>Prevalence of ADA Impact of additional DMARD therapy on ADA prevalence</i>	<i>Impact on clinical efficacy</i>	<i>Impact on side-effects</i>
B cell depletion (Rituximab and biosimilars)						
Strand et al.,2017 [28]	Systematic review	RA Patient demographics n/a	Data not available	0-21%	Patients with ADAs vs RTX showed less improvement in disease activity and were less likely to achieve clinical responses in RA patients. No comparative statistics/meta-analysis performed.	Higher rates of Tx emergent adverse events (89% vs 68%) were reported in patients with RA who develop anti-RTX ADAs compared to those who did not
Thurlings et al.,2010 [80]	The Netherlands Open-label cohort study	RA N=58 (F:M = 44:14)	Data not available	Data not available	Response to treatment and re-treatment measured by decrease in DAS28 and EULAR response was similar in ADA-positive and ADA-negative patients: p=0.87 and p=0.32 for the responses at 24 weeks after courses 1 and 2, respectively)	Data not available
Combier et al.,2020 [79]	France Retrospective cohort study	RA N=124 (F:M=97:27) Age (mean = 62; range 22-89) Other ARDS (including pSS, SLE, myositis) N=75	RA 13 years (1-60) Other ARDS 10 years (1-28)	RA 2.4% Other ARDS 14.7%	No data available on ADA impact on clinical efficacy 14.29% were tested because of loss of efficacy, and 78.6% were tested because of adverse reactions. No comparative statistics performed.	78.57% of ADA+ve patients (48/62 tested) with RA and other ARDs had infusion reactions to second or subsequent RTX cycles

		(F:M=59:16) Age (mean=57; range 21-85)				
Co-stimulatory blockade (Abatacept)						
Strand et al., 2017 [28]	Systematic review	RA (age 35-64) JIA (age 3-14.2) RA N = 1993 JIA N = Not available	RA: 1-54 JIA: 1-5	RA 2-20% JIA 2-11% Suggested that IV therapy associated with less immunogenicity than SC	Data not available	Data not available
Doeleman et al., 2019 [39]	Systematic review and meta- analysis	JIA IV N=190 SC N=173 Mean age IV – 12.4 (3.0) SC – 13.0 (10.0- 15.0)	IV – 4.4 (3.8) SC – 2.0 (0.0-4.0)	9.9% (pooled from 3 studies) (95% CI = 0.3–28.6)	No association between ADA and treatment failure was found	No injection site reactions experienced with SC and no adverse reactions for IV formulations were described
Hara et al., 2019 [76]	Japan Open label, multicentre single arm study	JIA IV N=20 Mean age 10.5 years (5-16) 4-8 years – 40% 9-12 years – 35% 13-17 years – 25%	0.75 (0.2-11.9)	5% (IV only)	No association between immunogenicity and loss of efficacy was found No comparative statistics performed	No association with safety, adverse events or hypersensitivity was found.
Brunner et al., 2018 [74]	International open label, multicentre	JIA N=219	2-5 years, 0.5 (0.0-1.0)	2.3% - 6-17 years 8.7% 2-5 years (SC only)	No clinical significance of ADA was found.	No issues regarding safety were found.

	study single arm study	2-5 years, N=46, median age – 4.0 (3.0-5.0) 6-17 years, N=173, median age – 13.0 (10.0-15.0)	6-17 years 2.0 (0.0-4.0)			
Lovell et al., 2015 [77]	Multicentre RCT	JIA N=58 (active arm) N= 59 (placebo) Mean age 12.4± 2.9	3.8±3.8	Whole Abatacept molecule 3.4% (2/58) CTLA-4 region only 5.5% (9/58) (IV only)	No loss of efficacy was found in the two patients with anti-abatacept antibodies to the whole molecules. Of the 9 patients with ADA against the CTLA-4 region, 3 discontinued due to lack of efficacy (small sample size, so no comparative statistics performed).	No infusion reactions were experienced.
Haggerty et al., 2007 [73]	Integrated analysis across multiple double blind and open-label studies	RA N=2237	Data not available	RA 2.1% ADA+ve with MTX 2.3% vs ADA+ve without MTX 1.4% - not significant	Patients who discontinued had a higher level of ADA compared to those who did not discontinue (7.4% vs 2.6%). No comparative statistics performed	No adverse safety outcomes were described
IL-6 blockade (Tocilizumab/Sarilumab)						
Benucci et al.,2016 [81]	Italy Cohort study of Tocilizumab	RA N=126 (F:M = 110:16) Mean Age: 59±12 years Range: 26-83 years	Mean disease duration: 11±5 years	0.79% (1/126 patients)	The occurrence of ADA against Tocilizumab is very rare.	Data not available

Sigaux et al.,2017 [82]	France Cohort study of Tocilizumab	RA N=40 (F:M = 32:8) Mean Age: 56.5±14 years	16±11.7 months	3.2%	No association between ADA status and disease activity was found	
Burmester et al.,2017 [83]	Meta-analysis of phase III RCTs of Tocilizumab	RA TCZ-SC: N=3099 TCZ-IV: N=5875	Data not available	TCZ-SC: 1.5% TCZ-IV: 1.2%	No association with decreased clinical efficacy was found	No clear impact of ADA on safety and side effects was found
Yokota et al.,2014 [84]	Japan Phase II-III RCTs of Tocilizumab	sJIA N=67 (F:M = 38:29) Mean Age: 8.3±4.3 years	4.4±3.5 years	7.5%	No decrease in clinical effectiveness was reported	4/5 patients with ADA experienced mild to moderate infusion reactions
Burmester et al.,2017 [85]	Multicentre RCT of Sarilumab	RA N=184 (F:M = 157:27) Mean Age: 50.9±12.6 years	8.1±8.1 years	7.1%	ADA were not associated with a loss of efficacy	ADA were not associated with hypersensitivity reactions
Wells et al.,2019 [86]	USA Open label study of Sarilumab	RA N=132 (F:M = 106:26) Mean Age: 52.4±13.4 years	10.5±9.0 years	150mg: 12.3% 200mg: 6.1%	Persistent ADA were associated with lower sarilumab levels but no correlation with clinical efficacy	There was no evidence that ADA status was linked to adverse effects. No notable differences in hypersensitivity reactions based on ADA status (no comparative statistics performed)
Genovese et al.,2015 [87]	Multicentre RCT of Sarilumab	RA 150mg: N=400 50.1±11.9 years	150mg: mean 9.5 years (range: 0.3-44.7) 200mg:	150mg: 16.7% 200mg: 13.0%	The presence of ADA was not associated with discontinuations due to lack of efficacy.	The presence of ADA was not associated with hypersensitivity reactions

		200mg: N=399 50.8±11.8 years	8.6 years (0.3-34.2)			
Xu et al., 2019 [89]	Worldwide Two-compartment model study of Sarilumab	RA N=1770 (F:M = 1466:304) Mean Age: 52±12 years	Data not available	18%	ADA may be linked to higher drug clearance, but this study did not evaluate the impact on clinical efficacy	Data not available
IL-17 blockade (Secukinumab/Ixekizumab)						
Deodhar et al., 2019 [93]	Pooled clinical trial safety data for Secukinumab	PsA N=1380 (F:M = 742:638) Mean Age: 48.8±12.0 years AS N=794 (F:M = 265:529) Mean Age: 42.4±12.3 years	Data not available	<1% across all studies	No effect of ADA positivity on clinical efficacy was reported	Immunogenicity was not related to adverse effects
Mease et al., 2017 [94]	Multicentre phase III RCT of Ixekizumab	PsA N=417 (F:M = 225:192) Mean Age: 49.5±11.9	6.7±7.2 years	5.3%	72.7% (8/11) of ADA-positive patients achieved a clinical response. No comparative statistics performed as very small sample size	Data not available
Gordon et al., 2016 [95]	Combined phase III RCTs of Ixekizumab	Plaque psoriasis N=1150	Data not available	9%	19 patients (1.7%) with high titres of ADAs had a lower clinical response than that of patients with no or low-moderate ADAs (no p-value given).	Data not available

IL-12/23 blockade (Ustekinumab)						
Strand et al., 2017 [28]	Systematic review	PsA Patient demographics data not available	Data not available	8-11% Concomitant use of MTX, AZA, leflunomide or mycophenolate is associated with lower rates of ADAs against INF in PsA	Data not available	Data not available
Smolen et al., 2017 [90]	Multicentre RCT	RA 90mg/8wk N=55 (F:M=46:9) Age 50.8±13.0 RA 90mg/12wk N=55 (F:M=47:8) Age 51.1±10.6	RA 90mg/8wk 5.6 ±5.5 RA 90mg/12wk 6.8±5.9	RA: 5.7% (3.3% neutralising)	Data not available	Data not available
IL-1 blockade (Anakinra, Canakinumab and Rilonacept)						
Fleischmann et al., 2006 [96]	Multicentre RCT of Anakinra	RA N=1340 (F:M = 1045:354) Mean Age: 55.2 years (range: 19-85)	10.3 years (range: 0.2-59.5 years)	50.1% (1.9% neutralising)	52% of those with neutralising ADA reported disease progression (no comparative statistics performed)	No associations between ADA and adverse effects
Cohen et al., 2002 [97]	Multicentre RCT of Anakinra	RA N=419 Anakinra dose: 0.04mg/kg/day N=63 Mean Age: 52.6 years	0.04mg/kg/day: 6.3 years 0.1mg/kg/day 8.8 years 0.4mg/kg/day 7.0 years	2.7% (8 out of 297 screened for antibodies)	No impact on clinical efficacy was found	87.5% of ADA positive patients experienced injection site reactions. No p-value reported

		<p>0.1mg/kg/day N=74 Mean Age: 53.0 years</p> <p>0.4mg/kg/day N=77 Mean Age: 52.8 years</p> <p>1.0mg/kg/day N=59 Mean Age: 49.0 years</p> <p>2.0mg/kg/day N=72 Mean Age: 54.1 years</p>	<p>1.0mg/kg/day 6.5 years</p> <p>2.0mg/kg/day 8.0 years</p>			
Ilowite et al., 2009 [98]	Multicentre RCT of Anakinra	<p>JIA N=25 (F:M = 17:8)</p> <p>Mean Age: 10 years (range: 3-17)</p>	Mean: 3.9 years (range: 1-11)	72% (none were neutralising)	No impact on clinical efficacy was found	Data not available
Sun et al., 2016 [99]	Prospective Study of Canakinumab	<p>JIA N=201</p> <p>Age range: 2 to <20 years</p>		3.1% (6 of the 14 patients screened for antibodies were positive, giving an incidence of 6/196)	No evidence of loss in clinical efficacy was found. Observed trough canakinumab concentrations in ADA+ve patients were comparable to ADA- patients (no comparative statistics performed).	No association was demonstrated between ADA and adverse effects

Ruperto et al., 2012 [100]	Multicentre RCT of Canakinumab	JIA N=50 (F:M=28:22) Median Age: 8.0 years (IQR: 6.0-12.0)	Median: 2.7 years (IQR: 1.3-6.2)	8% (4/50 patients) None were neutralising.	Data not available	Data not available
Lovell et al., 2013 [101]	USA RCT of Rilonacept	JIA N=24 (F:M=16:8) Mean Age: 12.6±4.3 years	3.1 years (mean)	54.2% (13/24)	No correlation between ADA and clinical responses was found. Statistical testing not performed due to small sample size.	All patients who experienced ≥3 injection-site reactions were ADA-positive

Table 2- Impact of ADA on disease outcomes in children and adults with inflammatory arthritis treated with other biologic agents.

Legend: ARDS – autoimmune rheumatic diseases; AS – ankylosing spondylitis; JIA-juvenile idiopathic arthritis; PsA- psoriatic arthritis; pSS – primary Sjögren’s syndrome; RA- rheumatoid arthritis; RCT-randomised control trial; SLE – systemic lupus erythematosus.

<i>Prevalence of ADA</i>	<i>Adults with inflammatory arthritis</i>	<i>Children with juvenile idiopathic arthritis</i>
TNF-α blockers		
Adalimumab and biosimilars	0-67%	6-45%
Infliximab and biosimilars	6.1-62%	26-37%
Etanercept and biosimilars	0-13%	0-33%
Golimumab	2-39.9%	46.8%
Certolizumab	2.8-65%	Data not available
B cell depletion		
Rituximab and biosimilars	0-21%	Data not available
Co-stimulatory blockade		
Abatacept IV	2-20%	2-11%
Abatacept SC	2-20%	2-11%
IL-6 blockade		
Tocilizumab	0-16%	1-8%
Sarilumab	7-24.6%	Data not available
IL-17 blockade		
Sekukinumab	0-1%	Data not available
Ixekizumab	5.3-9%	Data not available
IL-12/23 blockade		
Ustekinumab	5.7-11%	Data not available
IL-1 blockade		
Anakinra	50.1-70.9%	81.8%
Canakinumab	Data not available	3.1-8%
Rinolcept	Data not available	54.2%

Table 3. Comparison between the prevalence ranges for ADA to various biologic agents in adult versus paediatric populations

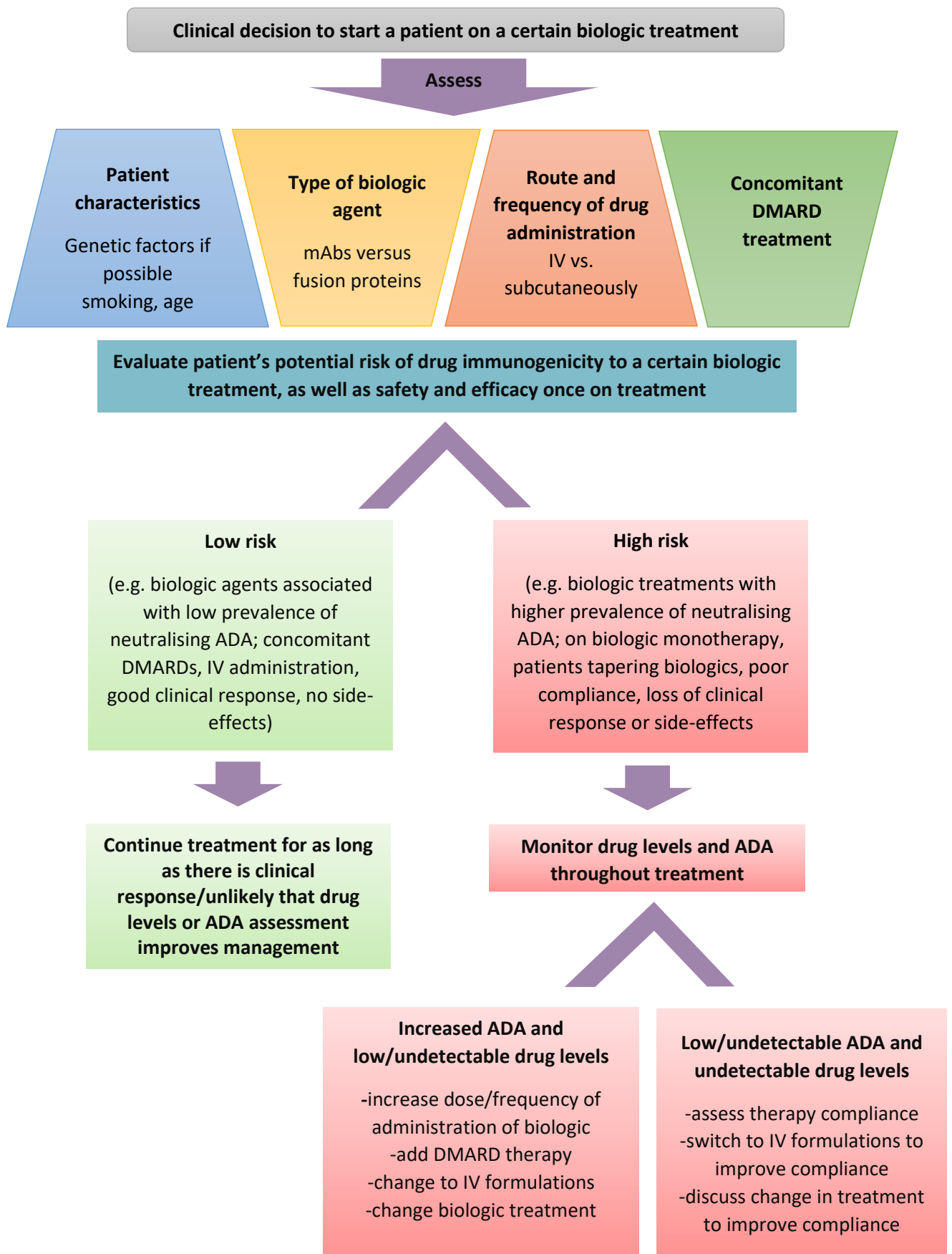


Figure 1: Potential clinical applications of the assessment of immunogenicity to biologic treatments