Sex-dimorphic adverse drug reactions to immune suppressive agents in inflammatory bowel disease

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Abstract

AIM: To analyze sex differences in adverse drug reactions (ADR) to the immune suppressive medication in inflammatory bowel disease (IBD) patients.

METHODS: All IBD patients attending the IBD outpatient clinic of a referral hospital were identified through the electronic diagnosis registration system. The electronic medical records of IBD patients were reviewed and the files of those patients who have used immune suppressive therapy for IBD, i.e., thiopurines, methotrexate, cyclosporine, tacrolimus and anti-tumor necrosis factor agents (anti-TNF); infliximab (IFX), adalimumab (ADA) and/or certolizumab, were further analyzed. The reported ADR to immune suppressive drugs were noted. The general definition of ADR used in clinical practice comprised the occurrence of the ADR in the temporal relationship with its disappearance upon discontinuation of the medication. Patients for whom the required information on drug use and ADR was not available in the electronic medical record and patients with only one registered contact and no further follow-up at the outpatient clinic were excluded. The difference in the incidence and type of ADR between male and female IBD patients were analyzed statistically by $\chi^2$ test.

RESULTS: In total, 1009 IBD patients were identified in the electronic diagnosis registration system. Out of these 1009 patients, 843 patients were eligible for further analysis. There were 386 males (46%), mean age 42 years (range: 16-87 years) with a mean duration of the disease of 14 years (range: 0-54 years); 578 patients with Crohn’s disease, 244 with ulcerative colitis and 21 with unclassified colitis. Seventy percent (586 pts) of patients used any kind of immune suppressive agents at a certain point of the disease course, the majority of the patients (546 pts, 65%) used thiopurines, 176 pts (21%) methotrexate, 46 pts (5%) cyclosporine and one patient tacrolimus. One third (240 pts, 28%) of patients were treated with anti-TNF, the majority of patients (227 pts, 27%) used IFX, 99 (12%) used ADA and five patients certolizumab. There were no differences between male and female patients in the use of immune suppressive agents. With regards to ADR, no differences between males and females were observed in the incidence of ADR to thiopurines, methotrexate and cyclosporine. Among 77 pts who developed ADR to thiopurines, 54 pts (65%) used thiopurines, 176 pts (21%) methotrexate, 46 pts (5%) cyclosporine and one patient tacrolimus. One third (240 pts, 28%) of patients were treated with anti-TNF, the majority of patients (227 pts, 27%) used IFX, 99 (12%) used ADA and five patients certolizumab. There were no differences between male and female patients in the use of immune suppressive agents. With regards to ADR, no differences between males and females were observed in the incidence of ADR to thiopurines, methotrexate and cyclosporine. Among 77 pts who developed ADR to one or more anti-TNF agents, significantly more females (54 pts, 39% of all anti-TNF treated women) than males (23 pts, 23% of all anti-TNF treated men) experienced ADR to an anti-TNF agent [$P = 0.011$; odds ratio (OR) 2.2, 95%CI 1.2-3.8]. The most frequent ADR to both anti-TNF agents, IFX and ADA, were allergic reactions (15% of all IFX users and 7% of all patients treated with ADA) and for both agents a significantly higher rate of allergic reactions in females compared with males was observed. As a result of ADR, 36 patients (15% of all patients using anti-TNF) stopped the treatment, with significantly...
higher stopping rate among females (27 females, 19% vs 9 males, 9%, \( P = 0.024 \)).

**CONCLUSION:** Treatment with anti-TNF antibodies is accompanied by sexual dimorphic profile of ADR with female patients being more at risk for allergic reactions and subsequent discontinuation of the treatment.

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**Key words:** Adverse drug reactions; Sexual dimorphism; Infliximab; Adalimumab; Inflammatory bowel disease

**INTRODUCTION**

The existence of a sex dimorphic profile of adverse drug reactions (ADR) has been increasingly recognized in recent years. Several studies on various therapeutics pointed to differences between sexes in the incidence as well as character of ADR. In general, females seem to be more at risk for ADR to various medication, 70% of drug users with ADR in a large cohort of 2367 patients being women[9]. In addition to this generally increased risk of ADR, female patients also differ from males in terms of types of ADR to a range of medication such as antiarthritics, antipsychotics, anti-retroviral drugs, and analgesics[2,3]. A limited number of small size studies performed in the field of auto-immune diseases and transplant medicine suggested existence of a sexual dimorphic profile of ADR to immunsuppressive medication, but this has not been studied in depth. Male sex has been reported as a risk factor for nodular regenerative hyperplasia in inflammatory bowel disease (IBD) patients treated with azathioprine[6]. In rheumatoid arthritis patients’ population, males have been shown to be more at risk than females for methotrexate-associated interstitial pneumonia[5] and to bacterial pneumonia complicating treatment with infliximab (IFX)[9]. For females, a higher incidence of azathioprine-related alopecia has been reported in transplant recipients[7]. In a pediatric population of Crohn’s disease (CD) patients, female sex was one of the risk factors for infusion reactions to anti-tumor necrosis factor (anti-TNF) treatment[8] and in an adult population of ankylosing spondylitis patients, females were more at risk of discontinuation of anti-TNF agents[9].

The sexual dimorphism of the immune responses is a generally accepted concept that has been studied predominantly in the context of the female predominance in autoimmune disorders[10] with the most important factor determining this dimorphism being the immunomodulatory properties of sex hormones. Considering these differences between the two sexes in basic immune reactions, further modulation of immune response by the immunosuppressive medication might have sex-specific consequences, including the quantitative and qualitative differences in ADR to these agents.

The limitations resulting from ADR for further therapeutic strategy in patients with chronic inflammatory conditions such as IBD are important. The ADR occurring in up to 20% of IBD patients using immune suppressive and anti-TNF agents[11,12] represent an important factor leading to the modulation or discontinuation of effective treatment. The thorough understanding of the underlying mechanism of ADR to these drugs, including the sex-related differences would help optimizing the therapy in these patients.

Therefore, in the present study we aimed to specifically determine the difference between male and female IBD patients in the occurrence and type of ADR to commonly used immunosuppressive agents, including ‘classical’ immune suppressive medication, i.e., thiopurines and methotrexate, as well as anti-TNF agents.

**MATERIALS AND METHODS**

**Patients**

All IBD patients attending the outpatient clinic of the Department of Gastroenterology and Hepatology of the Erasmus MC as of January 2009 were identified through the electronic diagnosis registration system. The medical records were reviewed with emphasis on details of drug treatment. Reported ADR to immune suppressive medication used for IBD were noted. Patients for whom the required information on drug use and ADR was not available in the electronic medical record and patients with only one registered contact and no further follow-up at the outpatient clinic were excluded.

**Definition of ADR**

All ADRs designated as such by the treating physician in the medical record were registered. The general definition of ADR used in clinical practice comprised the occurrence of the ADR in the temporal relationship with its disappearance upon discontinuation of the medication. In case of doubt about other concomitant factors contributing to the ADR a positive re-challenge was considered to be necessary for the event to be definitely categorized as ADR.

The ADR to immunosuppressive agents were divided into the following categories according to the type of symptom/event: gastro-intestinal, arthralgia and/or myalgia, cutaneous, infectious, malignancy, myelosuppression, hepatotoxicity, or pancreatitis. In case of anti-TNF
agents, additional categories of allergic reactions, lupus-like syndrome, and injection-site reactions were used.

Gastro-intestinal ADR comprised abdominal pain recognized by the patient as different from the IBD-related pain, diarrhea, nausea, and vomiting. Cutaneous ADR were defined as any kind of reported skin abnormality that occurred in temporal relationship with the treatment and resolved after cessation of the medication. Remittent or opportunistic infections occurring during the immunosuppressive treatment were noted as infectious ADR. For malignancies, any malignancy that was revealed during the treatment was categorized as ADR.

Myelosuppression was defined as leucopenia (leucocytes count < 4.0 × 10^9/L), and/or anemia (hemoglobin level < 8.5 mmol/L for males and < 7.5 mmol/L for females) and/or thrombocytopenia (thrombocytes count < 1.5 × 10^9/L). To categorize abnormal liver tests as hepatotoxic ADR, the increase of liver tests above 2 times upper normal value and absence of other causes, i.e., viral or autoimmune were required. Drug-induced pancreatitis were defined as any kind of reported skin abnormality occurring during the treatment.

Any of the following symptoms occurring during or within one day after infusion alone or in combination were considered as allergic reactions: skin reactions, dyspnoea, chest pain, low blood pressure, angioedema, fever and/or chills. Dyspnoea, skin abnormalities and arthralgia/myalgia occurring later than two days after infusion were categorized separately as potential delayed allergic reactions. Any motoric or sensoric loss, paresthesia and/or seizures were categorized as neurological ADR. Lupus-like syndrome diagnosis was characterized as the combination of arthritis and/or flu-like symptoms or fever and presence of anti-nuclear and or anti-double strand DNA antibodies. Injection site reactions (applicable for adalimumab-ADA) were defined as pain or local skin reaction after injection.

Non-specific ADR which could not be categorized according to these criteria were analyzed together and are further referred as others.

### Demographic characteristics and the use of medication

In total, 1009 IBD patients were identified in the electronic diagnosis registration system. Out of these 1009 patients, 843 patients were eligible for analysis according to the exclusion criteria stated in the methods part. There were 386 males (46%), mean age 42 years (range: 16-87 years) with a mean duration of the disease of 14 years (range: 0-54 years); 578 patients with CD, 244 with ulcerative colitis and 21 with unclassified colitis. There were no differences between male and female patients with regard to age and disease duration; significantly more males suffered from ulcerative colitis (141 pts, 58% of all ulcerative colitis patients).

Seventy percent (586 pts) of patients used any kind of immunosuppressive agents during the disease course, the majority of the patients (546 pts, 65%) used thiopurines, 176 pts (21%) methotrexate, 46 pts (5%) cyclosporine and one patient tacrolimus. No differences between male and female patients were observed with regard to the frequency of use of immunosuppressive agents in general or of particular agent.

One third (240 pts, 28%) of patients were treated with anti-TNF; the majority of patients (227 pts, 27%) used IFX, 99 (12%) ADA and five patients certolizumab. There were no sex-related differences in the use of anti-TNF agents (Table 1).

### Statistical analysis

The sex-related differences in categorical variable were analyzed statistically using two-sided χ² testing, for continuous variables a two-sided independent t-test was used. P-values < 0.05 were considered significant. The analysis was performed using SPSS PASW 17 software.

### RESULTS

#### ADR to immune suppressive agents

In total 278 patients experienced ADR to immunosuppressive agents; of which 155 patients to thiopurines (28% of all thiopurine-treated patients), 44 to methotrexate (25%), 2 (4%) to cyclosporine, and 77 pts (32%) developed ADR to one or more anti-TNF agents. As the age may represent an important factor influencing the development of ADR, a subanalysis was performed to compare the mean age between the groups of patients with and without ADR to immune suppressive agents and to anti-TNF agents. The mean age between the patients with and without ADR did not differ neither in the groups with ADR to immune suppressive agents (mean ± SEM, 41 ± 1 years in the group with ADR vs 40 ± 0.7 years in the group without ADR; P = 0.76) nor in the group with ADR to anti-TNF (mean ± SEM, 37 ± 1.5 years in the group with ADR vs 40 ± 1 year in the group

### Table 1 Demographic characteristics and the use of medication (n = 843)

|                | Males¹ n (%) | Females¹ n (%) | P value² |
|----------------|--------------|----------------|---------|
| Males/females  | 386 (46)     | 457            |         |
| Age (mean, range, yr) | 43 (16-87)    | 42 (18-87)     | 0.138   |
| Duration of the disease (mean, range) | 14 (0-48)      | 14 (0-54)      | 0.168   |
| CD/UC/unclassified | 233 (40)/141 (58) | 345/103/9 < 0.0001 |
| Immunosuppressive agents | | | |
| Thiopurines | 247 (45)    | 299 (46)       | 0.665   |
| Methotrexate | 73 (42)     | 103 (45)       | 0.203   |
| Cyclosporine | 26 (57)     | 20 (46)        | 0.170   |
| Tacrolimus  | 0           | 1              |         |
| Anti-TNF     | 101 (42)    | 139 (42)       | 0.109   |
| Infliximab   | 93 (41)     | 134 (46)       | 0.102   |
| Adalimumab   | 39 (41)     | 60 (47)        | 0.328   |
| Certolizumab | 1 (20)      | 4 (47)         | 0.401   |

1For each categorical variable number and percentage of males within the group is displayed; ²P values for test of the sex-related differences; two-sided χ² test for categorical variables and t-test for continuous variables.

Anti-TNF: Anti-tumor necrosis factor agents; CD: Crohn’s disease; UC: Ulcerative colitis.
patients experienced an ADR, one developed pseudomembranous colitis following treatment but she was also treated with systemic steroids. The second patient had a cutaneous reaction to cyclosporine.

Among 77 pts who developed ADR to one or more anti-TNF agents, significantly more females (54 pts, 39% of all anti-TNF treated women) than males (23 pts, 23% of all anti-TNF treated men) experienced ADR to an anti-TNF agent \( P = 0.011; \) odds ratio (OR) 2.2, 95%CI 1.2-3.8 (Figure 1C). In the subanalysis of respective anti-TNF agents, significantly more females (51 pts, 38% of all IFX-treated women) than males (20 pts, 22% of all IFX-treated men) suffered from ADR to IFX \( P = 0.009; \) OR 2.2, 95%CI 1.2-4.1) (Figure 1D). Relatively more females than males experienced ADR to ADA (16 females; 28% vs 6 males; 15%), this difference was not significant though \( P = 0.216 \) Figure 1D.

The most frequent ADR to both IFX and ADA were allergic reactions (15% of all IFX users and 7% of all patients treated with ADA) and for both agents a significantly higher rate of allergic reactions in females com-

Figure 1 Relative percentages of male and female patients experiencing adverse drug reactions to “classical” immune suppressive agents (thiopurines, methotrexate, cyclosporine and tacrolimus) and anti-tumor necrosis factor agents. A: Adverse drug reactions (ADR) to immune suppressive agents in general; B: ADR to respective immunosuppressive agents; C: ADR to anti-tumor necrosis factor agents (anti-TNF) in general; D: ADR to respective anti-TNF agents. Two-sided \( \chi^2 \) test. Black bars: ADR present; Gray bars: No ADR.

without ADR to anti-TNF; \( P = 0.79 \).

Overall, there were no significant differences in the frequencies of the ADR to immune suppressive agents between males and females. In total, 27% of males (71 pts) experienced ADR to any kind of immune suppressive agent compared to 34% of females (108 pts; \( P = 0.087 \)), (Figure 1A).

Among thiopurines users, 26% of males (63 pts) and 31% of females (92 pts) suffered from ADR to thiopurines \( P = 0.183 \); Figure 1B). The most frequent ADR to thiopurines were myelosuppression, hepatotoxicity (both in 33 pts, 6%) and gastro-intestinal ADR (23 pts, 4%), (Table 2). No differences between male and female patients were observed with regard to specific type of ADR to thiopurines.

In the group of patients treated with methotrexate, 27% (20 pts) of males and 23% (24 pts) of females experienced ADR \( P = 0.597 \); Figure 1B). The most frequent ADR to methotrexate were hepatotoxicity and gastro-intestinal ADR (both in 10 pts, 6%) (Table 2).

Out of 46 pts treated with cyclosporine, two female patients experienced an ADR, one developed pseudomembranous colitis following treatment but she was also treated with systemic steroids. The second patient had a cutaneous reaction to cyclosporine.

Among 77 pts who developed ADR to one or more anti-TNF agents, significantly more females (54 pts, 39% of all anti-TNF treated women) than males (23 pts, 23% of all anti-TNF treated men) experienced ADR to an anti-TNF agent \( P = 0.011; \) odds ratio (OR) 2.2, 95%CI 1.2-3.8 (Figure 1C). In the subanalysis of respective anti-TNF agents, significantly more females (51 pts, 38% of all IFX-treated women) than males (20 pts, 22% of all IFX-treated men) suffered from ADR to IFX \( P = 0.009; \) OR 2.2, 95%CI 1.2-4.1) (Figure 1D). Relatively more females than males experienced ADR to ADA (16 females; 28% vs 6 males; 15%), this difference was not significant though \( P = 0.216 \) Figure 1D.

The most frequent ADR to both IFX and ADA were allergic reactions (15% of all IFX users and 7% of all patients treated with ADA) and for both agents a significantly higher rate of allergic reactions in females com-
pared with males was observed (Table 3). There were no other sex-specific ADR observed to anti-TNF agents.

Of 77 patients experiencing ADR to an anti-TNF agent, 36 pts stopped the treatment (47%; overall discon- tinuation rate 15%), 27 pts (35%) switched to another anti-TNF agent, and 14 pts (18%) continued the treat- ment. As a result of ADR, 27 females (9% of all anti-TNF-treated males; 6 pts, 6% of all anti-TNF-treated women) stopped the treatment compared with males was observed (Table 3). There were no other sex-specific ADR observed to anti-TNF agents.

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### Table 2 Categories of adverse drug reactions to immunosuppressive agents in general and stratified by sex n (%)

| Category                        | Thiopurines | Thiopurines<sup>1,2</sup> | Methotrexate | Methotrexate<sup>1,2</sup> |
|---------------------------------|-------------|---------------------------|--------------|-----------------------------|
| Myelosuppression                | 33 (6)      | 12 (56)                   | NA           | NA                          |
| Hepatotoxicity                  | 33 (6)      | 19 (58)                   | 10 (6)       | 6 (60)                      |
| Pancreatitis                    | 10 (2)      | 2 (20)                    | NA           | NA                          |
| Gastro-intestinal side-effects  | 23 (4)      | 8 (35)                    | 10 (6)       | 6 (60)                      |
| Arthralgia and/or myalgia       | 11 (2)      | 5 (45)                    | 2 (1)        | 1 (50)                      |
| Cutaneous side-effects          | 10 (2)      | 4 (40)                    | 3 (2)        | 3 (100)                     |
| Infectious                      | 6 (1)       | 3 (50)                    | 2 (1)        | 1 (50)                      |
| Others<sup>3</sup>              | 23 (4)      | 7 (30)                    | 13 (7)       | 2 (13)                      |
| Not specified<sup>1</sup>       | 6 (1)       | 3 (50)                    | 4 (2)        | 1 (25)                      |

<sup>1</sup>No significant difference observed between males and females for none of the ADRs as tested with two-sided χ² test.  
<sup>2</sup>Number and percentage of males in the group of patients with adverse drug reactions (ADR) to thiopurines;  
<sup>3</sup>Number of all the people treated with the medicine;  
<sup>4</sup>Headache, paresthesia, hair loss, fatigue, emotional instability, malaise; specific for methotrexate-injection site reaction. NA: Not available.

### Table 3 Types of adverse drug reactions to respective anti-tumor necrosis factor agents in general and stratified by sex n (%)

| Category                        | Infliximab<sup>1</sup> | Infliximab<sup>1,2</sup> | P value<sup>2</sup> | Adalimumab | Adalimumab<sup>1,3</sup> | P value<sup>2</sup> |
|---------------------------------|------------------------|--------------------------|--------------------|-----------|--------------------------|--------------------|
| Allergic reaction               | 33 (15)                | 7 (21)                   | 0.045              | 7 (7)     | 0                        | 0.039              |
| Cutaneous SE                    | 9 (4)                  | 5 (56)                   | 0.492              | 3 (3)     | 3 (100)                  | 0.064              |
| Neurological SE                 | 4 (1.8)                | 0                        | 0.146              | 2 (2)     | 0                        | 0.512              |
| Dyspnoea                        | 4 (1.8)                | 0                        | 0.146              | 1 (1)     | 0                        | 1.00               |
| Arthralgia and/or myalgia       | 11 (5)                 | 4 (45)                   | 1.00               | 3 (3)     | 1 (33)                   | 1.00               |
| Injection site reactions<sup>1</sup> | NA                    | NA                       | 0.016              | 3 (3)     | 1 (33)                   | 1.00               |
| Infectious SE                   | 0                      | 0                        | 0.41               | 3 (3)     | 0                        | 1.00               |
| Malignancy                      | 1<sup>*</sup> (0.4)    | 16 (100)                 | 0.41               | 0         | 0                        | 0.0                |
| Lupus-like                      | 1 (0.4)                | 0                        | 0                  | 0         | 0                        | 0.0                |
| Others<sup>3</sup>              | 8 (3.5)                | 2 (25)                   | 0.476              | 0         | 0                        | 0.0                |

<sup>1</sup>Number of all the people treated with the medicine;  
<sup>2</sup>Two-sided χ² test to compare the differences in the frequencies of specific adverse drug reaction between males and females;  
<sup>3</sup>Number of males;  
<sup>*</sup>In case of adalimumab;  
<sup>3</sup>Abdominal pain, hair loss, fatigue, malaise, emotional instability;  
<sup>4</sup>Testis carcinoma. NA: Not available.

### DISCUSSION

In this large retrospective study, we studied sex differences in the frequency and types of ADR to immune suppressive medication in IBD patients. In contrast to thiopurines and methotrexate with similar rates of ADR in both sexes, we observed a sex dimorphic profile of ADR to anti-TNF agents with higher frequency of ADR among female IBD patients compared with male pa- tients. With regard to particular types of ADRs, females experienced more often allergic reactions to the most frequently used anti-TNF agents, IFX and ADA. In addi- tion, these ADR have led to discontinuation of treatment more frequently in females than males, thus substantially limiting the long term use of anti-TNF agents by female patients.

The landmark randomized controlled trials on IFX and ADA efficacy and safety did not reveal sex dimorphic profile of ADR to anti-TNF agents<sup>13-17</sup>. However, the design of these studies with rather short follow-up might underestimate the overall incidence of ADR and immune-mediated ADR occurring at long term in par- ticular. This would subsequently limit the sample size to study specific risk factors for the development of ADR.
Another source of information on ADR, the safety registry with inclusion of patients being at the physicians’ discretion are difficult to interpret with regard to sex differences in ADR incidence due to the possible selection bias and also a rather short follow-up. Interestingly, in a large retrospective, real-life study on long-term safety of IFX for the treatment of IBD, female sex was shown to be an independent risk factor for the development of hypersensitivity reactions and dermatological ADR to IFX. In addition, one small size study with pediatric CD patients determined female sex as one of the risk factors for infusion reactions. Thus, our results, in line with previous reports, suggest that female IBD patients are at specific risk for hypersensitivity reactions to monoclonal anti-TNF antibodies.

For each drug group with sex dimorphic ADR profile specific considerations for underlying mechanisms are applicable. The basic pharmacokinetic differences between the genders were at first considered to cause the predominance of ADR to some drugs in females, but over the past years, it became evident that sex hormones interacted with the particular drugs’ metabolism and mechanisms of action. The sex hormones greatly influence the immune responses which might also account for the sex differences in ADR to immune system modulating therapy. However, in the present study, only biological anti-TNF agents and no other immunosuppressive treatment showed a specific ADR sex dimorphic profile. In addition, the particular ADR presenting more frequently in females were hypersensitivity reactions, suggesting a female-specific immunogenic potential related to biological therapy.

ADR to monoclonal anti-TNF antibodies have been shown to be related to the development of antibodies against these agents. Anti-infliximab as well anti-adalimumab antibodies are found in the sera of patients with loss of response and/or ADR to IFX or ADA, suggesting thus the humoral immune response as underlying mechanism of IFX- and ADA- related immunogenicity. Interestingly, many studies analyzing the sex differences in the humoral response to vaccinations showed higher antibody titers in females and more local and systemic adverse reactions to influenza and rubella vaccines have been observed in women compared with men. The underlying mechanism of these sex differences in humoral immune response in general is not elucidated thus far, but taken all these observational data together, it is tempting to speculate that the use of monoclonal antibodies against TNF-α might result in higher anti-anti-TNF antibodies formation rate in female patients which in turn would lead to the female-specific higher incidence of immune-mediated ADR.

This immunogenic potential of anti-TNF agents resulting in ADR has important consequences for the management of IBD patients using these drugs. In this study, up to 50% of patients experiencing ADR discontinued the treatment, with the overall discontinuation rate of all patients using anti-TNF agents being 15%. There are thus considerable potential limitations of long term use of these otherwise effective drugs. Understanding the underlying mechanisms, one of which might be female-specific immunogenicity would subsequently help to identify patients at risk and modify the therapeutic strategy accordingly.

To our knowledge, this is the first report studying the sex differences in ADR profile to immunosuppressive medication in a large cohort of patients with immune-mediated disease. The main limitation of our study is its retrospective design in which reporting bias of ADR is inevitable due to the lack of a standardized protocol that would ensure a meticulous screening of every treated patient in a prospective study design.

In conclusion, female IBD patients are at increased risk of hypersensitivity reactions to anti-TNF agents compared with males. These ADR have important clinical consequences as they lead to the discontinuation of the treatment in half of the patients experiencing these reactions. Further research, with a specific consideration of sex dimorphism of the immunogenic potential of anti-TNF agents is warranted in order to improve the clinical management of patients at risk for ADR.

**COMMENTS**

**Background**

Adverse drug reactions (ADR) to immune suppressive agents used for inflammatory bowel disease (IBD) limit the treatment in up to 20% of patients. For various drugs, sex dimorphic profile of ADR has been documented but this has not been studied in depth in case of immune suppressive medication.

**Research frontiers**

Sex has great impact on the various physiological and pathophysiological processes including pharmacokinetics and pharmacodynamics. The immune responses differ between men and women and therefore, it is presumable that the response to immune suppressive treatment may have specific profile in male and female patients. This could specifically apply for the ADR that are immune-mediated, such as the reactions to biological therapy. With the current situation of immunogenicity of biologicals being the main limitation of this otherwise effective therapy, identifying patients with high immunogenic potential is one of the possible strategies to enhance the overall success of anti-tumor necrosis factor agents (anti-TNF) therapy.

**Innovations and breakthroughs**

In the present study, authors show that female IBD patients are more at risk for allergic reactions to infliximab and adalimumab compared with male patients. This sex-dimorphic profile of ADR seems to be specific to biological therapy and not a general feature of immune suppressive medication since no sex-specific ADR to thiopurines and methotrexate were observed. This increased risk for ADR to biological has been reported previously in small sample size studies of patients with various immune-mediated conditions but has not been studied as a particular phenomenon in IBD patients.

**Applications**

The described phenomenon of increased immunogenic potential of anti-TNF...
agents in females compared to males has important consequences for the management of IBD patients using these drugs. In this study, up to 50% of patients experiencing ADR to anti-TNF discontinued the treatment, with the overall discontinuation rate of all patients using anti-TNF agents being 15%. There are thus considerable potential limitations of long term use of these otherwise effective drugs. Understanding the underlying mechanisms, one of which might be female-specific immunogenicity would subsequently help to identify patients at risk and modify the therapeutic strategy accordingly.

Peer review
This retrospective study in a cohort of 843 patients is aimed to specifically determine the difference between male and female IBD patients in the occurrence and type of ADR related to immunosuppressive agents (thiopurines, methotrexate, cyclosporine, tacrolimus), and anti-TNF agents (infliximab, adalimumab, certolizumab). The results indicate that IBD women are at increased risk for allergic reactions in the course of anti-TNF treatment.

REFERENCES

1. Tran C, Knowles SR, Liu BA, Shear NH. Gender differences in adverse drug reactions. J Clin Pharmacol 1998; 38: 1003-1009
2. Francioni F, Brunelleschi S, Stardo L, CUomo V. Gender differences in drug responses. Pharmaco 2007; 55: 81-95
3. Nicolson TJ, Mellor HR, Roberts RR. Gender differences in drug toxicity. Trends Pharmacol Sci 2010; 31: 108-114
4. Vernier-Massouille G, Cosnes J, Lemenn M, Martova P, Reinsch W, Lacharrre D, Cadiot G, Bourhiuk Y, Deos M, Bryelle A, Duclos B, Seksik P, Mary JY, Colombel JF. Nodular regenerative hyperplasia in patients with inflammatory bowel disease treated with azathioprine. Gut 2007; 56: 1404-1409
5. Shidara K, Hoshi D, Inoue Y, Yamada T, Nakajima A, Taniguchi A, Hara M, Momohara Sar, Kamatani N, Yamanaka H. Incidence of and risk factors for interstitial pneumonia in patients with rheumatoid arthritis in a large Japanese observational cohort, IORRA. Mod Rheumatol 2010; 20: 280-286
6. Takeuchi T, Tatsuki Y, Nogami Y, Ishiguro N, Tanaka Y, Yamanaka H, Kamatani N, Hanigai M, Ryu J, Inoue K, Kondo H, Inokuma S, Ochi T, Koike T. Immunogenicity of two versus three injections of adalimumab for maintenance therapy of Crohn's disease: the ACCENT I randomised trial. Lancet 2006; 368: 1541-1549
7. Hanauer SB, Sandborn WJ, Rutgeerts P, Fedorak RN, Lukas M, MacIntosh D, Panaccione R, Wolf D, Pollack P. Human anti-tumor necrosis factor monoclonal antibody (adalimumab) in Crohn's disease: the CLASSIC-I trial. Gastroenterology 2006; 130: 1235-1247quiz 391
8. Rutgeerts P, Sandborn WJ, Feagan BG, Reinsch W, Olson A, Johanns J, Travers S, Rachmilewitz D, Hanauer SB, Lichtenstein GR, de Villiers WJ, Present D, Sands BE, Colombel JF. Infliximab for induction and maintenance therapy for ulcerative colitis. N Engl J Med 2005; 353: 2462-2476
9. Sandborn WJ, Hanauer SB, Rutgeerts P, Fedorak RN, Lukas M, MacIntosh D, Panaccione R, Wolf D, Kent JD, Bittle B, Li J, Pollack PF. Adalimumab for maintenance treatment of Crohn's disease: results of the CLASSIC II trial. Gut 2007; 56: 1232-1239
10. Sands BE, Anderson FH, Bernstein CN, Chey WY, Feagan BG, Fedorak RN, Kamm MA, Korzenik JR, Lashner BA, Onken JE, Rachmilewitz D, Rutgeerts P, Wild G, Wolf DC, Marsters PA, Travas SB, Blank MA, van Deventer SJ. Infliximab maintenance therapy for fistulizing Crohn's disease. N Engl J Med 2004; 350: 876-885
11. Lichtenstein GR, Feagan BG, Cohen RD, Salzberg BA, Diamond RH, Chen DM, Pritchard ML, Sandborn WJ. Serious infections and mortality in association with therapies for Crohn's disease: TREAT registry. Clin Gastroenterol Hepatol 2006; 4: 621-630
12. Bartelds GM, Wijbrands CA, Nurmoohamed MT, Stapel S, Lens BF, Aarden L, Dijkmans BA, Tak PP, Wolbink GJ. Clinical response to adalimumab: relationship to anti-adalimumab antibodies and serum adalimumab concentrations in rheumatoid arthritis. Ann Rheum Dis 2007; 66: 921-926
13. Pascual-Salcedo D, Plascencia C, Ramiro S, Nuño L, Bonilla G, Nagore D, Ruiz Del Agua A, Martinez A, Aarden L, Martin-Mola E, Balsa A. Influence of immunogenicity on the efficacy of long-term treatment with infliximab in rheumatoid arthritis. Rheumatology (Oxford) 2011; 50: 1445-1452
14. Radstake TR, Svensson M, Ejsbout AM, van den Hoogen FH, Enevold C, van Riel PL, Bendtzen K. Formation of antibodies against infliximab and adalimumab strongly correlates with functional drug levels and clinical responses in rheumatoid arthritis. Ann Rheum Dis 2009; 68: 1739-1745
15. Vultaggio A, Matucci A, Nencini F, Pratesi S, Parronchi P, Rossi O, Romagnani S, Maggi E. Anti-infliximab IgE and non-IgE antibodies and induction of infusion-related severe anaphylactic reactions. Allergy 2010; 65: 657-661
16. West RL, Zelinkova Z, Hub E, Rutgeerts P. Inflammation and drug toxicity. 2010; 69: 130-142
17. Giacometti A, Majani S, Nogami Y, Ishiguro N, Tanaka Y, Yamanaka H, Kamatani N, Hanigai M, Ryu J, Inoue K, Kondo H, Inokuma S, Ochi T, Koike T. Formation of antibodies against infliximab and adalimumab strongly correlates with functional drug levels and clinical responses in rheumatoid arthritis. Ann Rheum Dis 2009; 68: 1739-1745
18. Vultaggio A, Matucci A, Nencini F, Pratesi S, Parronchi P, Rossi O, Romagnani S, Maggi E. Anti-infliximab IgE and non-IgE antibodies and induction of infusion-related severe anaphylactic reactions. Allergy 2010; 65: 657-661
19. West RL, Zelinkova Z, Wolfink GJ, Kuipers EJ, Stokkers PC, van der Woude CJ. Immunogenicity negatively influences the outcome of adalimumab treatment in Crohn’s disease. Aliment Pharmacol Ther 2008; 28: 1122-1126
20. Chen XQ, Bilbiliar D, de Gast GC, van Loon AM, Nalin DR, van Hattum J. Immunogenicity of two versus three injections of inactivated hepatitis A vaccine in adults. J Hepatol 1997; 26: 260-264
21. Chiaramonte M, Majari S, Ngatchu T, Moschen ME, Baldove R, Zennisli G, Simonello L, Rocco S, Bertin T, Naccarato R, Trivello R. Two different dosages of yeast derived recombinant human adalimumab. J Hepatol 2002; 37: 373-378
22. Gower et al. Sex-dimorphic adverse drug reactions to anti-TNF alpha monoclonal antibodies in Crohn's disease. N Engl J Med 2004; 350: 876-885
23. Zhang DN, van den Hoogen FH, Enevold C, van Riel PL, Bendtzen K. Formation of antibodies against infliximab and adalimumab strongly correlates with functional drug levels and clinical responses in rheumatoid arthritis. Ann Rheum Dis 2009; 68: 1739-1745
24. Chen XQ, Bilbiliar D, de Gast GC, van Loon AM, Nalin DR, van Hattum J. Immunogenicity of two versus three injections of inactivated hepatitis A vaccine in adults. J Hepatol 1997; 26: 260-264
25. Chiaramonte M, Majari S, Ngatchu T, Moschen ME, Baldove R, Zennisli G, Simonello L, Rocco S, Bertin T, Naccarato R, Trivello R. Two different dosages of yeast derived recombinant human adalimumab: a comparison of immunogenicity. Vaccine 1996; 14: 135-137
26. Benjamin CM, Chew GC, Silman AJ. Joint and limb symptoms in children after immunisation with measles, mumps, and rubella vaccine. BMJ 1992; 304: 1075-1078
27. Gower et al. Sex-dimorphic adverse drug reactions to anti-TNF alpha monoclonal antibodies in Crohn's disease. N Engl J Med 2004; 350: 876-885
28. S-Editor Gu SX L-Editor A E-Editor Zhang DN