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Discontinuation of infliximab therapy in patients with Crohn’s disease in sustained complete remission (the STOP IT study): protocol for a double-blind, randomised, placebo-controlled, multicentre trial

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ABSTRACT

Introduction: Infliximab (IFX), a monoclonal chimeric antibody against tumour necrosis factor (TNF) α, is effective for induction and maintenance of remission in moderate to severe Crohn’s disease. Discontinuation of IFX maintenance therapy in patients in remission should be considered in order to reduce the potential long-term side effects and lower costs.

Methods and analysis: This is a prospective, double-blind, randomised, placebo-controlled, multicentre study of patients with luminal Crohn’s disease who have been treated with IFX for at least 1 year and are in sustained complete clinical, biochemical and endoscopic remission (ie, Crohn’s Disease Activity Index (CDAI) score <150, complete mucosal healing and biochemical markers of inflammation within the normal range). These patients are randomised to receive placebo infusions or continue IFX maintenance therapy. The primary end point is the proportion of patients in maintained remission after 48 weeks (def. CDAI <150).

Ethics and dissemination: It is estimated that the knowledge gained about how to optimally handle patients with Crohn’s disease in complete long-term sustained remission on IFX is proportionate to the risks and inconveniences related to participation in this study. Prolonged exposure to IFX may cause severe side effects and increased risk of malignancies. Conversely, IFX discontinuation should not unnecessarily create a high risk of relapse. Thus, empirical evidence is needed concerning the safety of discontinuing IFX once a patient exhibits sustained remission. Study results will be published in an English language scientific medical journal. The study is approved by the Danish Medicines Agency (EudraCT-number: 2012-002702-51) and the Regional Ethics Committee of Region Hovedstaden Denmark (Approval-number: H-4-2012-099). The project is reported to the Danish Data Protection Agency (ID-number: 2007-58-0015/HEH.750.89-27), registered at Clinicaltrials.gov, and monitored by independent GCP units for the University of Copenhagen, Odense and Aarhus. The current approved protocol is V.3.2, dated 1 June 2014.

Trial registration number: http://clinicaltrials.gov/show/NCT01817426.

INTRODUCTION

Biological agents targeting tumour necrosis factor (TNF) α are effective in inducing and maintaining remission in patients with moderate-to-severe luminal and fistulising Crohn’s disease.1–4 The chronic nature of Crohn’s disease necessitates long-term TNF-inhibiting (TNFi) maintenance treatment. However, the potentially severe side effects such as infections, infusion reactions and increased risk of neoplasia,5 along with the high economic cost of the treatment, warrant the exploration of strategies for

Strengths and limitations of this study

▪ A state-of-the-art double-blind, randomised, placebo-controlled clinical trial. Previous studies are of retrospective origin, or post hoc analyses of prospective trials. Only two prospective observational studies have been done and these were performed without control groups.

▪ All evaluations are done by clinically validated scoring systems according to clinical trial practice (eg, Crohn’s Disease Activity Index, CDAI).

▪ Data analyses are not blinded.

▪ The selected patient population is narrow; consequently, the results can only be extrapolated to a fraction of patients on IFX maintenance therapy.
discontinuing TNFi among subgroups of patients in long-term sustained remission.

It is generally accepted that TNFi should not be discontinued in patients without complete remission. However, recent international guidelines conclude that existing data are insufficient to make firm recommendations on when and in whom to stop TNFi treatment. In contrast, several European countries, for example, UK and DK, now recommend re-evaluation of disease activity after 1 year of TNFi therapy to determine if the treatment should be discontinued.

The discontinuation decision is typically made from an individual judgement of benefits versus risks. Concerns related to the discontinuation of TNFi include the risk of disease flare, the risk of infusion reactions at re-initiation, and the possibility of reduced future medical treatment options. However, early treatment becomes more common and the long-run safety of TNFi has been questioned and its use is therefore not recommended any longer.16 However, data from the STORI study of 121 patients with luminal Crohn’s disease who discontinued IFX while in clinical remission after nearly 2 years (median 680 (412–948) days) have shown that 61% of patients with Crohn’s disease, who discontinued IFX while in complete remission, maintained remission for 1 year; indeed, half of the patients remained in remission after 1 year of TNFi therapy to determine if the treatment should be discontinued.

Recent prospective observational data from the STORI study of 115 patients with luminal Crohn’s disease found that 56% of patients maintained remission 1 year after discontinuation of infliximab (IFX). In a Hungarian prospective observational study of 121 patients with clinical remission, 45% had relapse within 1 year after cessation. Several retrospective studies reported similar relapse rates of 55–85%. Consistent with these data, we have reported that 61% of patients with Crohn’s disease, who discontinued IFX while in steroid-free IFX-induced remission, maintained remission for 1 year; indeed, half of the patients remained in remission after nearly 2 years (median 680 (412–948) days). Together with the current clinical experience, together with clinical remission follow the natural history of Crohn’s disease; prior effective treatment with IFX therefore does not appear to impose a subsequent disease-modifying effect. It has been suspected that the response to retreatment with IFX in case of relapse after IFX withdrawal may be lost. However, data from the STORI study and from our centre suggest that patients may respond well to retreatment with IFX at relapse.

The term ‘remission’ is not well defined and may incorporate one or more features such as clinical remission, as assessed by Crohn’s Disease Activity Index (CDAI) score, biochemical remission, endoscopic remission, etc. Patients who respond to TNFi therapy, clinical, biochemical and endoscopic, are considered to be in complete remission (typically defined as CDAI score <150 and no other signs of disease activity). Thus, the STORI study identified predictors of relapse, including certain features as well as objective biochemical and endoscopic markers of disease activity. The following risk factors were associated with relapse: (1) male sex; (2) absence of surgical resection; (3) corticosteroid use between 6 and 12 months before discontinuation of IFX; (4) IFX trough level ≥2 µg/mL at time of discontinuation of IFX; (5) Crohn’s Disease Endoscopic Index Score (CDEIS) >0; (6) leucocyte counts >6.0×10^9/L; (7) C reactive protein (CRP) ≥5.0 mg/L; (8) haemoglobin ≤145 g/L and (9) faecal calprotectin ≥300 µg/g. A complete model and a simplified model describing the risk of relapse with respect to the number of risk factors present were proposed. Interestingly, patients with ≤3 risk factors in the complete model, and ≤2 risk factors in the simplified model, had a very low risk of relapse of approximately 10% at 1 year after IFX discontinuation, as compared to a relapse rate of approximately 60% in those patients with a higher number of risk factors. Thus, degrees of remission might predict the outcome after discontinuation of IFX. Similar to these low-risk patients, relapse rates among patients with ongoing IFX therapy are 13% per year. Serum concentrations of TNFi in individual patients vary despite the same dosing. At initiation of treatment, high concentrations of TNFi immediately prior to the next administrations, that is, trough levels of the drug, are associated with maintenance of clinical remission. In addition, repeated infusions/injections of TNFi might result in the patient forming an antibody against the drug (anti-TNFAb). Development of anti-IFX Ab is generally associated with loss of response. However, patients in complete remission, with high levels of anti-TNFAb and low levels of TNFi, are functionally low drug exposure (low levels of TNFi with or without high levels of anti-TNFAb), might be able to tolerate cessation. Thus, monitoring patients in remission for circulating levels of functional TNFi as well as for anti-TNFAb is warranted to allow optimal individual treatment. Owing to methodological limitations, for example, cross-binding of drug and antidrug Ab in commonly used binding assays such as ELISA, levels are conventionally measured as trough levels. However, it is highly likely that this is not the optimal time point for assessments of rational therapeutic management.

As described above, there is a need to investigate how to manage patients with Crohn’s disease in long-term sustained complete remission. To date, there is little evidence whether IFX can be discontinued favourably and safely in this selected patient group. In prior studies, patients in remission who discontinued IFX therapy faced a higher risk of relapse than patients who continued therapy. However, data suggest that patients with very few risk factors (ie, in complete remission) might have almost similar relapse rates as if IFX therapy were maintained. The intention of this study is to examine whether IFX can be discontinued safely in the complete-remission patient group. To form a rational and efficient management algorithm, we will seek predictors of outcomes after discontinuation. Further, we
hypothesise that incorporation of measurements of blood levels of IFX and anti-IFX Ab is clinically relevant. A prospective controlled trial is suitable to investigate the optimal management in individual patients and to establish recommendation regarding withdrawal of IFX.

METHODS AND ANALYSIS

Study design and patients

This study is a prospective, double-blind (patient and physician), randomised, placebo-controlled, multicentre study. A study algorithm is shown in figure 1.

Adult patients (age ≥18 years) with luminal Crohn’s disease defined by standardised diagnostic criteria in sustained complete remission on IFX maintenance therapy (def. infusions every 6–10 weeks, 5–10 mg/kg) with a treatment length of a minimum 12 months are eligible. Complete remission is defined as a CDAI score <15017, no signs of inflammation on biochemical parameters (normal CRP, leucocytes, haemoglobin (Hb) and albumin), and no other marks of disease activity, either from endoscopic examination (Simple Endoscopic Score for Crohn’s Disease (SES-CD) score 0–2)24 25 and/or by MRI (defined by no signs of disease activity when evaluated by a trained radiologist).26 27 Sustained remission is defined as a clinical judgement of the disease to be stable at two consecutive treatment visits (corresponding two scheduled IFX infusions) and no use of oral steroids within 3 months prior to inclusion. Concomitant therapy with immune suppressants, except steroids, is allowed. The dosage and frequency must have been stable 3 months prior to inclusion, and must remain stable throughout the study period.

Exclusion criteria include the initial indication for IFX being predominantly fistulising perianal disease, ongoing fistulising disease, and pregnancy or lactation. Further, in case of any contraindications for continuing IFX treatment, including prior acute or delayed infusion reaction to a TNF-inhibiting agent, former malignancy, moderate to severe heart disease, any active infection requiring parenteral or oral antibiotic treatment, known infection with tuberculosis, HIV or hepatitis virus, the patient cannot be included.

Course of study

The course of this study is given in table 1 together with the study algorithm in figure 1. The total study length is 48 weeks.

The screening visit is defined as clinical remission and 12 months of scheduled IFX therapy minus the interval of regular visits. At the screening visit, biochemical parameters are assessed. Evaluation of mucosal healing is planned before the next visit; endoscopy for patients with colic/ileocolic Crohn’s disease; MRI for patients with small bowel Crohn’s disease and for patients with fistulising Crohn’s disease. The assessed biochemical parameters are: haemoglobin, leucocyte count, platelets, CRP, creatinine, alanine aminotransferase (ALAT), bilirubin, albumin and faecal calprotectin.

The inclusion visit is defined as the time of the next scheduled IFX infusion after the screening visit, thus resulting in an inclusion date after a minimum 12 months of IFX maintenance therapy.

All patients will be graded for disease activity in accordance with CDAI,28 Work Productivity and Activity

Figure 1 Study algorithm.
endoscopy (Simple Endoscopic Score for Crohn’s disease (SES-CD)) and/or MRI, and biochemical parameters at the time of inclusion. At the inclusion visit, patients are randomised (block randomisation and an allocation ratio of 1:1) to either continue IFX therapy at an unchanged dosage, or alternatively to receive matching placebo. Randomisation is stratified according to concomitant immunosuppressive therapy and history of fistulising disease. Randomisation is done centrally. The allocation sequence is in opaque, sealed envelopes. A non-blinded nurse, who is not involved in the treatment of the patient, receives the allocation result and subsequently prepares and labels IFX or placebo medication accordingly. Patients, treating nurses and treating physicians are blinded as to the type of medication (IFX or placebo) which the patient receives.

Following the screening and inclusion visit, patients are seen after 4 weeks; thereafter, the next consultations will be undertaken as part of the regular visits related to control, every 8 weeks. Disease activity will be assessed at every study-related visit by CDAI, WPAI, IBDQ and by biochemical parameters.

The study is terminated after 48 weeks (visit 9). At the time of visit 9, all patients will be graded for disease activity in accordance with CDAI, WPAI, IBDQ short, biochemical parameters and by endoscopic examination (SES-CD) and/or MRI. The treating physician and the patient will, after the termination of the study, be informed about whether the patient has continued IFX or received a placebo.

Patients who relapse during the course of the study will be withdrawn and treated at the discretion of the treating physician. It is the intention that patients who were randomised to discontinue IFX and subsequently relapse will be offered inclusion in an open label study of the effect of retreatment with IFX. This is, however, a separate study, not a part of the present study.

End points

The primary end point of this study is the proportion of patients who maintain remission at the end of the trial after 48 weeks. Remission is defined as CDAI<150.

Secondary end points assessed at the end of the trial after 48 weeks include:

- Proportion of patients who maintain complete remission (def. CDAI score <150, no signs of inflammation on biochemical parameters and mucosal healing).
- Median time to relapse after discontinuation of IFX.
- Proportion of patients experiencing relapse. Relapse, defined as a CDAI score >150 and a greater than 70 point increase from inclusion over two consecutive weeks.9,1
- Change from baseline in disease activity (ie, CDAI score, biochemical parameter including f-calprotectin, SES-CD).
- Economical expenses for treatment of Crohn’s disease in the two groups.

Sample size

Calculations are based on the assumption that remission rates are higher in patients maintaining IFX treatment compared to patients receiving a placebo. Relapse rates in patients during ongoing IFX therapy was estimated to be 13% per year in a recent review by Gisbert and Panés.18 As we will include a selected group of patients who have already received treatment for a year with good response, we expect the remission rate in patients

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**Table 1** Course of study and data collection in the intervention and control groups

<table>
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<th>Visit number</th>
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*Defined as time of next scheduled treatment.
†Acquire a signed written informed consent prior to conducting any study-related procedures.
‡Endoscopy and/or MRI between visits 1 and 2, and at 4–0 weeks before visit 9.
§C reactive protein, haemoglobin, white cell counts, platelets, albumin, creatinine, alanine aminotransferase, bilirubin and faecal calprotectin.
¶Blood sample for determination of infliximab (IFX) and anti-IFX antibody.
**Note, normally scheduled IFX therapy, not study medication.

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Index (WPAI),29 life quality score (IBDQ short),30,31 endoscopy (Simple Endoscopic Score for Crohn’s disease (SES-CD)) and/or MRI, and biochemical parameters at the time of inclusion. At the inclusion visit, patients are randomised (block randomisation and an allocation ratio of 1:1) to either continue IFX therapy at an unchanged dosage, or alternatively to receive matching placebo. Randomisation is stratified according to concomitant immunosuppressive therapy and history of fistulising disease. Randomisation is done centrally. The allocation sequence is in opaque, sealed envelopes. A non-blinded nurse, who is not involved in the treatment of the patient, receives the allocation result and subsequently prepares and labels IFX or placebo medication accordingly. Patients, treating nurses and treating physicians are blinded as to the type of medication (IFX or placebo) which the patient receives.

Following the screening and inclusion visit, patients are seen after 4 weeks; thereafter, the next consultations will be undertaken as part of the regular visits related to control, every 8 weeks. Disease activity will be assessed at every study-related visit by CDAI, WPAI, IBDQ and by biochemical parameters.

The study is terminated after 48 weeks (visit 9). At the time of visit 9, all patients will be graded for disease activity in accordance with CDAI, WPAI, IBDQ short, biochemical parameters and by endoscopic examination (SES-CD) and/or MRI. The treating physician and the patient will, after the termination of the study, be informed about whether the patient has continued IFX or received a placebo.

Patients who relapse during the course of the study will be withdrawn and treated at the discretion of the treating physician. It is the intention that patients who were randomised to discontinue IFX and subsequently relapse will be offered inclusion in an open label study of the effect of retreatment with IFX. This is, however, a separate study, not a part of the present study.

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who continue IFX maintenance therapy to be 90%. In
the STORI study, the subgroup with the best prognostic
markers had a similar proportion of patients maintain-
ing remission. However, it is unknown if this can be
extrapolated to other populations, especially because no
control group was included in the STORI study.12 Thus,
remission rates on discontinuation may be somewhat
lower. A difference between patients continuing and
patients stopping IFX of 20 percentage points is consid-
ered clinically relevant. Based on this minimally clinic-
ically relevant difference of 20% and an α set at 0.05 (two
sided) and a β set at 0.2, (IBM SPSS Sample Power V.3),
a total of 62 patients in each group are needed to dem-
onstrate a clinically relevant difference of continuing
versus discontinuing IFX therapy. To correct for drop-
outs, it is planned to include 136 patients in the study.

**Data analysis plan**

The inclusion period extends from 1 November 2012
until 31 October 2015. The study completion is sched-
uled for November 2016. Thereafter, we will analyse the
data and present results for scientific publications. In
order to obtain adequate participant enrolment, the
study employs multiple centres. The study is currently
ongoing at five Danish sites and remains open for par-
ticipation by new centres. The list of study sites can be
viewed at clinical.trial.org.

*Statistical analysis plan:* Descriptive statistics will be calcu-
lated as percentages for discrete variables, and median
with IQR or mean with SEM for continuous variables, as
appropriate. Fisher’s exact or $\chi^2$ test, as appropriate, will
be used for univariate analysis of discrete variables.
Unpaired t test/paired t test or Mann-Whitney U test/
Wilcoxon signed-rank test will be used for univariate ana-
lysis of continuous variables, as appropriate. Time until
relapse will be estimated using survival statistics, that is,
the Kaplan-Meier method and log-rank test. Association
of demographical, clinical and biochemical variables with
relapse will be estimated using univariate and multivaria-
ble Cox proportional hazard regression analysis. p Values
are two sided, and p<0.05 is considered significant.

**Explorative analyses**

In order to investigate whether pharmacoinmunological
data can rationalise therapeutic management with
respect to continuation or discontinuation of IFX
therapy, patients will, on the day of infusion, have three
blood samples drawn: one just before infusion (trough),
one right after the infusion (obtained from the other
arm) (peak) and one an hour after infusion (C₁). Based
on disease activity after 1 year, patients are categorised as
either in complete remission or not in complete remis-
ion. Results of IFX (and anti-IFX Ab) trough, peak and
C₁ concentrations in the two groups (remitters vs non-
remitters) will be compared. Samples will be measured
by common solid and fluid phase assays for this purpose,
for example, Reporter Gene Assay (RGA).16 20 32 33

**ETHICS AND DISSEMINATION**

*Ethical and safety considerations*

It is essential to determine strategies for discontinuation of
IFX therapy in patients with Crohn’s disease. Cessation of IFX therapy must not provide a very high
risk of the disease to flare. On the other hand, the pos-
sible risk for development of lymphoma, including hepato-
splenetic T-cell lymphoma, which is often fatal, and
other malignancies cannot be excluded. In particular,
young patients and patients who have received pro-
longed treatment are at risk.34 35 In addition, though
seldom, severe side effects to IFX present a danger for
patient safety.

Patients participating in this study may potentially
benefit from the discontinuation if they are able to toler-
ate cessation. The cessation can continue after the study
period, as long as they experience low disease activity. In
case of relapse, after discontinuation of IFX, data from
our study14 and from the STORI study12 suggest that the
patient will respond very well to retreatment. Patients
who have been randomly selected to continue IFX
therapy are subjected to the same side effects as they
would have faced before entering the study. Participation
in the trial should provide new data and information to
benefit future patients.

No study-related activities will be performed until
informed consent has been obtained. Investigators are
responsible for obtaining written consent from each
patient. During the trial, the patients will be monitored
closely and every reasonable precaution will be taken to
ensure patients’ safety. At each visit, adverse events
(AEs) will be elicited, and in case of any AEs a descrip-
tion will be recorded including intensity and causal rela-
tionship to the study drug. The principal investigator
must inform the Danish Medicines Agency immediately
if suspected unexpected serious adverse reactions
(SUSARs) occur during the trial. Investigators must
immediately report all serious incidents to the principal
investigator. In case of a serious AE or SUSAR, the blind-
ing for the involved patient is ceased. Patients will be
informed if there are major changes to the trial.
Investigators will be informed if there are important
protocol modifications. The trial will be prematurely ter-
ninated if scientific findings from other research groups
eliminate the trial. The trial will also be disrupted if
there are serious, not yet known, side effects to the
drugs used. IFX is a registered drug and is administered
following the current instructions for administering the
drug. Facilities for resuscitation and intensive therapy
will be immediately at hand and patients should be
under observation during the infusion.

The project will provide new knowledge regarding
how to optimally handle patients with Crohn’s disease in
sustained remission on a TNFi, and will help develop
new therapeutic strategies for this patient group. The
study will provide information about the value of meas-
uring concentrations of TNFi and antibodies against
TNFi in an everyday clinical decision-making set-up. In

addition, anti-TNF treatment plays an important role in many inflammatory diseases (eg, rheumatoid arthritis, spondylarthritides and psoriasis). As is the case in patients with Crohn’s disease, discontinuation of TNFi in patients with other inflammatory diseases—after achievement of low disease activity—is important for reasons of safety and economy. Thus, the study results might inspire and expand similar research studies in other inflammatory diseases impacting patient safety of TNFi therapies.

Publication
It is the intention that any results, positive, negative or inconclusive, will be published in a relevant English language scientific journal/conference. SSB will be the first author. The last authors will be MA, KB, JB, OØT and CS, the order of which will depend on the specific contributions in each manuscript. The site investigators will be the coauthors.

Data deposition and curation
We will store biological material in the form of blood samples from the patients. The samples are stored for 10 years after the completion of the study, after which all samples and leftover material will be destroyed. Patient-identifiable information (eg, name) will be treated as strictly confidential and will not be made publicly available. All information about the patients, including non-clinical data, protocols, case report forms and verbal and written information, are protected under the act concerning the processing of personal data and the Danish health law. Investigators will have access to the final trial data set. The study will comply with the ICH-GCP (Good Clinical Practice) rules and be performed in accordance with the protocol and relevant regulatory requirements.

Contributors All authors have been involved in the conception and design of the study. All authors contributed to the refinement of the study protocol. SSB, MAA and CS performed the drafting of the protocol manuscript. JB, OØT and KB have been involved in the revision of the manuscript. All authors approve the final manuscript and agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work will be appropriately investigated and resolved. They also participated in the implementation of the study.

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Competing interests CS has served as a speaker for MSD and Abbvie and as a consultant for MSD and Takeda Pharmaceutical Company. JB has served as an advisory board member for Abbvie. OØT has served as a speaker and consultant for UCB and Zealand Pharma, speaker for MSA, and primary investigator for Agen, Biogen, Novo-Nordisk and Pfizer. KB has served as a speaker for Pfizer, Roche, Novo-Nordisk, Bristol-Meyers Squibb and Biomonitor and owns stocks in Novo-Nordisk and Biomonitor.

Patient consent Obtained.

Ethics approval The Danish Medicines Agency (EudraCT-number: 2012-002702-51), The Regional Ethics Committee of Region Hovedstaden Denmark (Approval-number: H-4-2012-099). The Danish Data Protection Agency (ID-number: 2007-58-0015/HEH.750.89-27).

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement The study is registered at Clinicaltrials.gov (http://clinicaltrials.gov/show/NCT01817426).

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