Conclusion  Relapse rates after withdrawal of a thiopurine are high, particularly for CD, and predicting this remains difficult. To help increase the power of this study, we are now expanding it across the UK.

Competing interests  None declared.

REFERENCE

OC-167  CALCULATING THE “MISSED OPPORTUNITY” OF THIOPURINE MONOTHERAPY OVERCOME WITH THIOPURINE AND ALLOPURINOL COMBINATION THERAPY

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Introduction  A proportion of patients preferentially methylate thiopurines, resulting in high levels of methylated metabolites and low levels of thioguanine nucleotides. This can result in hepatotoxicity and treatment non-response. Co-prescription of thiopurines (at 25%–50% of standard dose) with allopurinol, (which blocks xanthine oxidase) circumvents this problem, optimising metabolite profile and clinical outcome and additionally overcomes atypical side effects experienced on thiopurine monotherapy. Using data from a large cohort of patients receiving combination therapy, we aimed to establish what proportion of all patients starting thiopurines could be salvaged from treatment failure to 1yr remission by combination therapy.

Methods  Using data from a cohort of 109 patients recruited retrospectively, all receiving combination therapy in our clinic, 1yr clinical response rates were calculated by indication. Using data from a published prospective cohort and side effect rates from meta-analysis, we calculated the proportion of all patients starting thiopurines that could be salvaged from treatment failure to 1yr remission by combination therapy.

Results  10/17 (59%) of hyper-methylating non-responders to thiopurine monotherapy, 8/17 (47%) of those treated for atypical side effects and 11/15 (75%) switched for hepatotoxicity achieved remission at 1 year. Using these response rates, potential gain was calculated from a prospective cohort (n=207) from our centre. 60 patients discontinued thiopurine monotherapy due to non-specific side effects, eight due to hepatitis and 32 were hyper-methylating non-responders, a total of 100 patients with clear indications for combination treatment. Our results predict that 53/100 could have achieved 1-year remission, representing 26% of the original cohort. Using a more conservative published side effect rate of 10% (Prefontaine et al 2010, Cochrane Database of Systematic Reviews, CD000545) and excluding 2.8% due to side effects unsuitable for combination therapy (pancreatitis and myelotoxicity), 12% of all patients started on azathioprine could have their outcome on thiopurine therapy converted from treatment failure to 1-year remission by combination therapy.

Conclusion  12%–26% represents the “missed opportunity” of patients starting thiopurine monotherapy which can be realistically overcome by combination treatment with allopurinol, converting treatment failure to successful 1-year remission. Given that thiopurines remain a key part of most IBD treatment paradigms, this is an important opportunity for improved treatment outcomes in IBD.

Competing interests  None declared.

OC-168  TOWARDS INDIVIDUALISED RISK PREDICTION FOR CROHN’S DISEASE

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Introduction  Although the RR of developing Crohn’s disease (CD) is increased in first degree relatives (FDR) of probands by 13–27-fold, (1) it is difficult to apply this on an individual basis. Recently, risk models for complex disease using information from genome-wide association and epidemiological studies have been built. Potentially, such models could identify FDR at highest risk of CD, for screening and early intervention.

Aim  To generate a risk model for CD, and ultimately determine if such a model can predict presymptomatic or future development of CD.

Methods  CD patients were contacted to identify asymptomatic FDR aged 18–55 who wished to participate in research into risk prediction and presymptomatic diagnosis of CD. (Recruitment target: 500 FDR) DNA was extracted from saliva. Genetic analysis used the Immunochip (Illumina), and genotypes were determined for disease susceptibility loci established by CD genome-wide meta-analysis. (2) A multiplicative risk model was generated using risk prediction software (REGENT) (3) based on genotype and smoking status, and CD RR determined for each FDR.

Results  The first 100 FDR have been recruited. Demographics: 53% siblings, 36% offspring, 11% parents. Mean age 37, females 66%, smokers 25%, ex-smokers 33%. NOD2 genotype counts were not significantly different to their frequencies in the control population, but were significantly different to cases (15% heterozygous, 0% homozygous, p=0.64 for controls vs FDR, p=0.001 for cases vs FDR). CD RR ranged from 0.04 to 10.07 (median 0.96) and was significantly higher than that in the general population: 19% of FDR had a RR >2, compared to an expected proportion of 11% (p=0.02). Comparison of highest and lowest RR quartiles in FDR showed good separation of RR between the groups. More FDRs in the highest RR quartile had >2 CD-affected relatives compared with FDRs in the lowest RR quartile, but this was not significant (5 vs 1, p=0.19).

Conclusion  Increased RR of CD is confirmed in FDR compared to the general population. Risk stratification is possible, but follow-up is required to determine if such risk scores can be used to predict presymptomatic CD or future development of CD. Additionally, risk awareness may encourage smokers to quit. Once recruitment to the study is completed, FDR in the highest and lowest risk quartiles will proceed to further tests including faecal and serum biomarkers, and capsule endoscopy. All those recruited will undergo follow-up for 10 years.

Competing interests  None declared.

REFERENCES