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The Royal Liverpool and **NHS Broadgreen University Hospitals NHS Trust** 

# Implementation of genotype-guided dosing of warfarin to improve anticoagulation control

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## Background

- Warfarin is an effective, widely used anticoagulant, but dosing is challenging due to its narrow therapeutic index and large interpatient variability in requirements.
- Needs close monitoring of international normalised ratio (INR), with target range typically 2-3. Important to establish therapeutic dose (that which maintains INR in target range) as soon as possible to reduce risk of adverse events (bleeding; thrombosis) and number of clinic visits required for INR monitoring.
- Many clinical, demographic and genetic factors associated with dose requirements, with variants in CYP2C9 and VKORC1 genes having the largest influence.
- EU-PACT trial<sup>1</sup> demonstrated genotype-guided dosing approach (GGD), using point-of-care genetic testing, led to patients spending 7% more time in target INR range and achieving target range sooner.
- We undertook an implementation project to determine whether GGD could translate into routine clinical practice in the UK.

## Study design and follow-up

- Matched cohort design: 3 clinics using GGD approach (implementation group); 3 comparable clinics using standard approach (control group).
- Statistical power boosted by using routinely collected INR data from similar anticoagulation clinics using standard approach (dashboard data).
- Patients in implementation group genotyped using point-of-care assay and dosed according to GGD days 1-5; dosed according to usual clinic practice thereafter. • Patients in control groups dosed according to usual practice throughout. All followed-up for 12 weeks.
- Data collected on: demographics, INR measurements, dose changes, withdrawals, hospital admissions. •
- Patient and staff questionnaires completed at implementation group clinics to gain feedback on GGD approach. •

## Point-of-care genotyping assay and dose calculator

### Genotyping assay:

- Buccal swab to obtain DNA, no DNA extraction required.
- ParaDNA point-of-care genotyping platform, developed by LGC.  $\bullet$
- Results available in 45 minutes.

#### **Dose calculator:**

Computerised web-based calculator incorporating loading dose (days 1-3) and maintenance dose (days 4/5) algorithms previously tested in EU-PACT.<sup>1</sup>

### Outcomes

- **Primary outcome:** % time in target INR range during first 12 weeks
- Secondary outcomes:
  - $INR \ge 4$  in first week

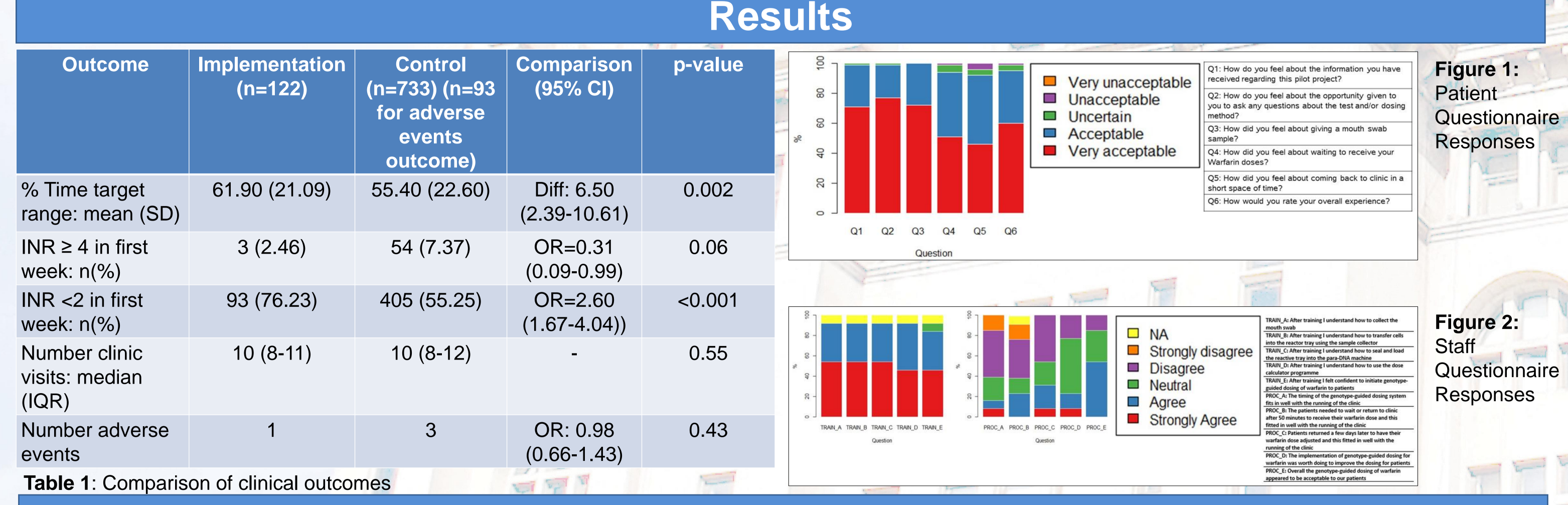
Patient adverse events (bleeds, mortality, other morbidity)

- INR <2 in first week
- Total number clinic visits in first 12 weeks

- 5. Patient opinion of GGD
- 6. Staff opinion of GGD

### **Statistical methods**

- Time in target INR range calculated using method of Rosendaal et al.<sup>2</sup>
- Student's t-test, chi-square test and Mann-Whitney U test used as appropriate to compare between implementation and control groups (with and without dashboard data, lacksquarewhere applicable). Significance threshold of 0.05 assumed.
- Descriptive analysis of questionnaire responses.



### Conclusions

Despite increasing popularity of direct acting oral anticoagulants (DOACs), warfarin remains the most cost-effective anticoagulant for a majority of patients. Further, DOACs are contraindicated for some patient subgroups including those with severe renal impairment, on certain interacting drugs and children. However, it is essential for effectiveness and patient safety that therapeutic dose of warfarin is achieved quickly and maintained. Results of our project demonstrate that the GGD approach supports this goal (Table 1), can be implemented smoothly into clinical practice with only a few minor modifications (Fig 2), and moreover is viewed positively by patients (Fig 1) and staff (Fig 2).

#### References

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2. Rosendaal FR, Cannegieter SC, van der Meer FJ, Briet E. A method to determine the optimal intensity of oral anticoagulant therapy. Thromb Haemost. 1993;69(3):236-9.