Opt-out bloodborne virus screening: a cross-sectional observational study in an acute medical unit

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ABSTRACT

Objective Recent treatment developments for HIV, hepatitis C virus (HCV) and hepatitis B virus (HBV) have greatly improved prognoses. Current screening practices are mainly risk based and are suboptimal. Improved efforts are critically needed to identify persons with these viruses. The aims of this study were to assess the feasibility of an opt-out bloodborne virus (BBV) screening programme in an acute medical unit (AMU) and to describe the prevalence of HIV, HBV and HCV in this population.

Design and setting This was a cross-sectional observational study in the AMU of a tertiary referral hospital in Galway, a city in the west of Ireland.

Participants 1936 patients entered the study; 54% were male, mean age was 53.1 years (SD 19.6). During the study period, all patients attending the AMU aged ≥16 years who were having bloods drawn and who had the ability to verbally consent for an additional blood sample met the inclusion criteria for the study.

Results Over 44 weeks, 1936/4793 (40.4%) patients consented to BBV panel testing. Diagnosed prevalence rates for HIV, HBV and HCV were 0.5/1000, 2/1000 and 1.5/1000, respectively. There was one HIV-positive result; the patient was already engaged in care. Four patients tested positive for HBV surface antigen; one new diagnosis, one previously lost to follow-up and two already engaged in care. Three patients had active HCV infection; two had been lost to follow-up and are now linked back into services.

Conclusion BBV testing uptake of 40.4% is higher than previous studies in AMU settings that used opt-in strategies, but lower than expected, possibly due to not incorporating testing into routine practice. The diagnosed prevalence of HBV is notable as little data currently exist about its prevalence in Ireland. These data are valuable in order to inform further prevention strategies for these infections in low-prevalence settings.

INTRODUCTION

Infection with HIV, hepatitis C virus (HCV) and hepatitis B virus (HBV) account for significant morbidity and mortality globally, including in Ireland; screening has both personal and public health benefits.1–3 All three viral infections can be diagnosed on serum samples, and for all three infections, treatment is available which leads to better patient outcomes and lower rates of onward transmission.4–5 Late diagnosis can lead to increased morbidity and mortality, increased use of resources and increased costs of care.12 Furthermore, these infections often occur in marginalised groups who are less likely to present for routine testing and therefore who are more likely to present late with advanced disease.3–6–8

HIV

Worldwide an estimated 0.8% (0.7%–0.9%) of adults aged 15–49 years are living with HIV,6 and in Europe there are between 30000 and 33000 new cases of HIV reported each year.10 Previous studies have estimated the diagnosed prevalence rates of HIV in Ireland (table 1).610 To date in Ireland, 5253 people have been diagnosed with HIV, but it is estimated that, in keeping with statistics for the rest of Europe, about 15% of HIV infections remain undiagnosed.1314 Approximately 90% of those with diagnosed HIV infection in Ireland are on treatment, and 90% of these
Table 1  Current screening practices and demographic information for BBVs in Ireland

| Current screening practice | Recommended screening practice | Prevalence | Main transmission modes | Population affected |
|----------------------------|--------------------------------|------------|-------------------------|--------------------|
| HIV Risk based STI clinics Antenatal screening Blood donation (opt-out) | Universal in areas of high prevalence (≥2/1000) 14| Estimated 1/1000 11 2.25/1000 in Dublin ED 8| Sexual injection drug use | MSM PWID |
| HIV Risk based Blood donation (opt-out) | Risk based Opt-out in areas of high prevalence 24 | 0.5%–1.2% 23 Estimated 60% of infections undiagnosed 24–28 | 80% PWID Recent increase in sexual transmission (MSM) 10 | PWID MSM—threefold rise since 2003 10 |
| HBV Risk based STI clinics Antenatal screening Blood donation | Screening for all immigrants coming from areas where HBV prevalence >2% 32 | <1% | Vertical transmission (occurring outside of Ireland) 7 | Persons from countries of high endemicity 7 |

**Table notes:**
- **BBV:** bloodborne virus; **ED:** emergency department; **HBV:** hepatitis B virus; **HCV:** hepatitis C virus; **MSM:** men who have sex with men; **PWID:** people who inject drugs; **STI:** sexually transmitted infection.

patients are virally suppressed, 10 suggesting that our main gap to achieving United Nations Programme on HIV/AIDS (UNAIDS) 90/90/90 targets is around diagnosis. 13 14

Active case finding of undiagnosed HIV infection leads to better prognosis for individual patients and prevents the onward transmission of HIV by viraemic patients who are unaware of their HIV status. 15 The Center for Disease Control (CDC) estimates that if everyone with HIV infection was diagnosed, there would be approximately one-third fewer new sexual HIV transmissions. 15

The Strategic Timing of Antiretroviral Therapy (START) trial (August 2015) showed clinical benefits for patients who started treatment early in their disease course, which requires diagnosing infection before it presents clinically, 16 and a recommendation for earlier treatment initiation has been incorporated into international guidelines. However, with our current HIV testing methods, patients are continuing to present late (defined as a CD4 cell count <350). Forty-five per cent of those presenting with HIV in Ireland in 2015 presented late. 3

Previous studies have shown that healthcare workers are also a barrier to testing. Testing has previously been associated with a laboured process of pretest counselling and written consent; many providers are unaware of the changes in testing guidelines, and importantly are unaware of the lack of need for written consent or extensive pretest counselling. 15 17 18

Current recommendations call for doctors to offer people HIV tests if they are at high risk of infection, 19 but many healthcare providers do not make this assessment. Furthermore, by adopting this targeted testing approach, patients with high-risk behaviours may feel stigmatised by being singled out for HIV testing. Opt-out testing among pregnant women has already proven to be highly effective in preventing mother to child transmission of HIV by picking up unsuspected cases of HIV. 20–22 CDC and UK National Guidelines both recommend opt-out HIV testing in patients presenting to healthcare services, but this strategy has not yet been widely implemented (table 1). 15 19

**Hepatitis C virus**

The prevalence of HCV in Ireland is estimated at 0.5%–1.2% (table 1). 23 It is estimated that there are between 20 100 and 42 000 people with current infection in Ireland, and that up to 60% of these infections remain undiagnosed. 24–28 Injection drug use is the most common risk factor associated with HCV acquisition in Ireland (80%), followed by possible sexual exposure (5%), receipt of blood or blood products (4%), vertical transmission (2%) and tattooing or body piercing (1%). 24 Of note, there have been an increase in sexually transmitted HCV infections nationally in men who have sex with men from 4 in 2014 to 29 in 2016. 25

Chronic undiagnosed HCV infection can lead to liver cirrhosis, hepatocellular carcinoma and death. 23 The recent advances in treatment with oral, well-tolerated shorter course curative treatment with direct-acting antivirals (DAAs) for HCV have resulted in cure rates of >90% in those treated. 23 The Health Service Executive (HSE) National Hepatitis C Strategy 2011–2014 made recommendations across four key areas, which included expanded screening and treatment, after the recent availability of DAAs for HCV in Ireland in 2014. 24

**Hepatitis B virus**

While Ireland has a low HBV prevalence (thought to be <1%), from 2002 to 2008 there were a large number of new notifications in persons from areas of high endemicity, such as central Asian republics, South East Asia and sub-Saharan Africa (>8% prevalence) (table 1). 7 8

The overall prevalence of HBV in the EU/EEA is low at 0.9% (range 0.1%–4.4%). 29 30 According to the European Centre for Diseases Control data, Ireland still has one of the highest rates of HBV and HCV new notifications in Europe. 30 Eighty per cent of notifications of chronic HBV
in Ireland in 2013 were in persons from a country of high HBV prevalence or in persons with asylum seeker status.7

HBV infection can be controlled with antiviral medication, preventing complications related to liver failure. Onward transmission of HBV occurs more frequently than other BBVs but can be prevented by viral suppression through treatment in infected individuals and through vaccination of close contacts.7 The US CDC guidelines recommend HBV testing in those with exposure risk factors.31 The 2015 Health Protection Surveillance Centre guidelines recommend screening all immigrants who are coming from countries where prevalence of HBV is >2%.32

### Diagnosis and screening

No widespread screening programme is in place for HIV, HCV and HBV. Current screening practices for HIV and HBV in Ireland are mostly risk based, although CDC and UK guidelines call for opt-out screening in certain circumstances, for example, blood donation (table 1).15 19 A recent study in Dublin showed high feasibility and acceptability of opt-out BBV screening in the emergency department (ED) of a tertiary referral hospital, with a cumulative uptake rate of 50.1%.6 Based on this, the study protocol has become routine practice in this ED, with ongoing opt-out BBV screening. A recent study in primary care centres in Dublin also showed high acceptability rate, with 89.5% uptake on opt-out BBV screening.33

The aims of this study were to assess the feasibility of an opt-out BBV screening programme in an acute medical unit (AMU) of a tertiary referral hospital in Galway, a city in the west of Ireland, and to describe the prevalence of HIV, HBV and HCV in this population. This study is, to our knowledge, the first to assess the feasibility of an opt-out screening approach for BBVs in an AMU in a low diagnosed prevalence area for BBVs. It is also the first to describe the prevalence of HIV, HBV and HCV in an AMU.

### METHODS

This was a cross-sectional observational study conducted in the AMU of Galway University Hospital in the city of Galway, Ireland. The study involved collaboration between the AMU and the departments of Infectious Diseases, Microbiology, Virology and Hepatology.

Patients are referred to the AMU from the ED, or directly from general practitioners. All patients are medically stable. At registration, each patient was given a patient information leaflet by the administration staff to review and ask any further questions about the study. At least 20 min later, and with verbal consent, blood was drawn by nursing staff. Each clinical proforma used for routine clinical care was stamped with a reminder for the provider to ask the patient about consent to have bloods drawn for the study. In line with international guidelines for HIV testing and with local ethical approval, verbal consent was deemed sufficient for all three BBVs.15 19 A streamlined process was developed by the microbiology/virology team to manage the testing for all three viruses within a service which provides over 5000 new patient consultations per year. The study duration and sample size were determined by funding available. Patients did not receive any reimbursement for their inclusion in the study.

All samples were tested routinely, using the Abbott HBsAg Qualitative II, Abbott anti-HCV and Abbott HIV Ag/Ab Combo assays. Results were processed in the on-site microbiology laboratory. Positive results were reported by phone to the study team as per routine clinical care. Results were managed on a ‘no news is good news’ policy, with a dedicated phone line for patients to ring if they wanted to follow-up results or had any questions relating to the study.

Inclusion criteria were patients presenting new to the AMU who were aged 16 years or older and were having bloods drawn for any reason and who had capacity to consent to inclusion in the study. Targets for uptake of screening were set at 50% of patients accepting to participate in the study for the first 2 months, and 80% from month 3 onwards, based on the Dublin ED study targets and results.6 Linkage to care was coordinated by the study team where appropriate. Data analysis was primarily descriptive and was done using SPSS V.24. Prevalence rates for HIV, HCV and HBV infections were described and the feasibility of the study was assessed by the uptake of BBV testing.

### RESULTS

From 18 January to 21 November 2016, 1936/4793 (40.4%) patients were assessed for medical care in the department consented to BBV panel testing. A total of 1941 samples were each tested for HIV, HBV and HCV (five duplicate samples). Fifty-four per cent of the patients were male and their mean age was 53.1 years (SD 19.6) (figure 1). The diagnosed prevalence of HIV was 0.5/1000. There was one HIV-positive result; this patient was previously diagnosed and engaged in care. The diagnosed prevalence of HBV was 2/1000. Four patients tested positive for HBV surface antigen; one new diagnosis, one previously lost to follow-up and now linked back into care and two already engaged in specialist care. One patient testing positive for HBV was >65 years of age. This patient came from a high-endemic area for HBV. The diagnosed prevalence of HCV was 1.5/1000. Six patients were HCV...
antibody (Ab) positive. Three patients had active HCV infection; two had been lost to follow-up and are now linked back into services, the third was already engaged in care. Of the other three patients that were found to have evidence of HCV Ab positivity, only one declined follow-up bloods to assess for active infection; they had a prior documented undetectable viral load. Fifty per cent (3/6) of the patients who tested positive for HCV Ab were >65 years of age.

The overall uptake of BBV screening over the course of the study was 40.4% (figure 2). The decline in testing from week 20 to week 23 reflects the unexpected relocation of the AMU department to a different section of the hospital. After this relocation, there was a further educational drive around testing; however, there was also a tail-off in testing towards the end of the study which affected the overall uptake (figure 2). One patient changed his/her mind about being included in the study, having initially agreed. There were no issues reported by patients to the study team either in person or via the dedicated phone line regarding the opt-out screening process.

DISCUSSION
The results of our study suggest that an AMU BBV screening programme is feasible. The percentage uptake of 40.4% is higher than previous studies in AMU settings that used opt-in strategies (6%–22%). This uptake was calculated based on the assumption that all patients attending the AMU had bloods drawn and therefore were eligible for the study. There may have been some patients who did not have bloods done, in which case this uptake would be a slight underestimate. However, the uptake of testing was lower than expected, and lower than the cumulative 50.1% uptake achieved in an ED setting in Dublin, the capital city of Ireland, which has a higher volume of patients, including more critically unwell patients. It was also significantly lower than the uptake of 89.5% achieved in a study conducted across four urban primary care sites in Dublin. This was a research study and we believe that the uptake in our study was affected by not incorporating testing into routine clinical care. The significantly lower uptake in the AMU and ED studies compared with the 89% uptake achieved in the primary care setting may reflect the difference between care by a single provider in the community versus a larger team of providers in an ED or AMU setting (table 2).

The overall diagnosed prevalence rate of HIV of 0.5/1000 was lower than previously reported. This likely reflects the possibility that patients triaged to AMU are not fully representative of the local population, tending to be older and often having more medical comorbidities. A similar study in the ED would be needed to ascertain the true prevalence of BBVs in the local population presenting to the hospital.

This prevalence of 0.5/1000 does not meet the 2/1000 suggested in the British recommendations for opt-out testing for HIV. However, the study is still in line with WHO guidelines released as early as 2006 that recommend all patients presenting to healthcare between the ages of 13 and 64 should be tested at least once for HIV. Furthermore, modelling in the USA suggests that routine screening for HIV infections is cost-effective and comparable to costs of other routinely offered screening where the prevalence of HIV exceeds 0.5/1000.

The diagnosed prevalence of HBV of 2/1000 is notable as little data currently exist around HBV prevalence in Ireland. The urban ED study found a HBV prevalence rate of 5/1000 and the urban primary care setting showed a similar prevalence to ours of 2/1000 (table 2). From a public health perspective, these data are valuable in order to inform further screening and prevention strategies for these infections in a low-prevalence setting. Understanding of rates of BBVs in different areas, and the patient demographics, is imperative when planning screening systems. In our case, all patients with HBV were from areas of high endemicity for HBV and therefore a risk-based screening programme should capture these infections.

We found a low HCV diagnosed prevalence rate of 1.5/1000, compared with 50/1000 in the Dublin ED setting. This was expected in light of the reduced risk factors in our catchment area compared with Dublin, the most notable of these is the low prevalence of people who inject drugs (PWID) in the local community. Despite this,
our prevalence results are in keeping with the estimated overall prevalence of HCV in Ireland, which is 0.5%–1.2%, and are only slightly lower than the prevalence of 2/1000 found in the urban primary care setting33 (table 2). As for HBV, little published data existed prior to this study about prevalence rates of HCV in our low-prevalence population, so these data are also valuable in informing further screening practices.

We feel that healthcare worker-associated barriers also contributed to our uptake being lower than expected, and lower than the uptake in the primary care study. This could be improved by education of healthcare practitioners about the lack of need for extensive pretest counselling and written consent, and by incorporating testing into routine care. Moreover, in areas of low prevalence targeted testing may be more difficult to achieve as people may not disclose their risk factors to their doctor. We believe that this study helped awareness about BBVs in the AMU and we feel that opt-out testing removed the stigma associated with BBV testing for both patient and the provider, as has been described previously.36

Notably 3/6 (50%) of the patients who tested positive for HCV Ab in our study were over 65 years of age and would not have met routine screening criteria based on WHO guidelines.15 One 69-year-old man was unaware of having previously been infected and clearing HCV, and did not have any risk factors that would have been picked up by targeted testing. One patient testing positive for HBV was >65 years of age. This patient came from a high-endemic area and could have been picked up if targeted screening was systematically in place for patients coming from areas of high endemicity; however, this is not currently the case. Ireland also has an ageing HIV population and older patients may be less likely to be picked up with current screening practices.11

Our study had a number of limitations. In designing the study, a decision was made about the importance of a streamlined consent process; the study needed to fit in with routine work activities. Due to this, we could not collect any patient feedback or comparative data between patients who did and did not participate in the study. The study design of including all adults attending the AMU aimed to eliminate any bias in patients being included in the study. However, there is the chance that those who were in fact at high risk for BBV (particularly HIV and HCV) may have declined screening, and therefore our study may have underestimated prevalence. Another limitation is that a small number of patients may not have had bloods drawn, in which case the uptake would be underestimated. Risk demography was not collected and was difficult to ascertain retrospectively but this limitation is similar in opt-in studies. An unexpected challenge was the relocation of the AMU during the study, which was not foreseen and which impacted all activities, including this study, as seen by the decline in uptake from week 20 to week 23.

**CONCLUSION**

The results of this study suggest that opt-out BBV screening is feasible in an acute medical assessment unit setting, in an area of low prevalence for BBVs. The method of opt-out testing removes the stigma of BBV testing for both the patient and the provider.36 The percentage uptake of 40.4% is higher than previous studies in AMU settings that used opt-in strategies.11 However, the uptake of testing was lower than expected, and lower than that which was achieved in an urban ED setting6; this was a research study and we believe the uptake was affected by not incorporating testing into routine clinical care. The study has added valuable information on prevalence of BBVs in the study population. The diagnosed prevalence of HBV is most notable as little data currently exist about its prevalence in Ireland. The results suggest that a widespread AMU BBV screening programme throughout Ireland may be feasible. Modelling studies have been done to try to assess the cost-effectiveness of screening for HIV in high-prevalence settings,35 but to the best of our knowledge, there is no published data on cost-effectiveness of simultaneous BBV screening either inside or outside of Ireland. Further research is required to determine the cost-effectiveness of such a strategy and to evaluate the incidence and prevalence of BBVs in other catchment areas and healthcare settings.

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**Table 2** Uptake rates and bloodborne virus prevalence rates across three study sites: urban primary care, urban emergency department and low-prevalence acute medicine unit

|          | Uptake  | HIV prevalence (/1000) | HBV prevalence (/1000) | HCV prevalence (/1000) |
|----------|---------|------------------------|------------------------|------------------------|
| Primary care | 89.5% | 0                      | 2                      | 2                       |
| Emergency department | 50.1% | 11                     | 50.5                   | 5                       |
| Acute medicine unit | 40.4% | 0.5                    | 2                      | 1.5                     |

HBV, hepatitis B virus; HCV, hepatitis C virus.
Contributors HT, DG, CB and SOC designed the study protocol. HT and DG gained ethical approval for the study. NA, CF, AD, AL, MS, MBK and NB participated in the data collection. NA, AL, MS, MBK, NB, JL and CF ensured patients were linked to appropriate clinical care where required. CF, EMC and DK supervised the processing of laboratory samples and collated the laboratory data. NA, HT and DG analysed and interpreted the collated data. NA wrote the manuscript and all the authors reviewed the manuscript prior to publication.

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Competing interests None declared.

Patient consent for publication Not required.

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