N_LyST: a simple and rapid screening test for Lynch Syndrome

Susanti Susanti1,2,3*, Wakkas Fadhil1,3*, Henry O. Ebili1,3,4*, Asiri Abutaleb1,3, Ausrine Nestarenkaite5, Efthymios Hadjimichael1,3 Hersh A. Ham-Karim1,3, Joanne Field6, Katherine Stafford6, Balawir Matharoo-Ball7, James C.Hassall1,3, Abid Sharif6, Anca Oniscu8, Mohammad Ilyas1,3

* Co-first authors

1Molecular Pathology Group, Unit of Academic Molecular Pathology, Division of Cancer and Stem Cell, School of Medicine, University of Nottingham, Queen’s Medical Centre Campus, The University of Nottingham, Nottingham, UK. 2Department of Pharmacology and Clinical Pharmacy, Faculty of Pharmacy, University of Muhammadiyah Purwokerto, Central Java, Indonesia. 3Nottingham Molecular Pathology Node, University of Nottingham, Queen’s Medical Centre Campus, The University of Nottingham, Nottingham, UK. 4Department of Morbid Anatomy and Histopathology, Olabisi Onabanjo University, Ago-Iwoye, Nigeria. 5National Center of Pathology, Affiliate of Vilnius University Hospital Santaros Klinikos, Lithuania. 6East Midlands Regional Molecular Genetics Service, Nottingham University Hospitals NHS Trust, Nottingham, UK. 7Nottingham Health Sciences Biobank, Nottingham University Hospitals NHS Trust, Nottingham, UK. 8Molecular Pathology, Royal Infirmary of Edinburgh, 51 Little France Crescent, Edinburgh. EH16 4SA.

Corresponding author: Prof M Ilyas

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ABSTRACT

Background and aims
We sought to use PCR followed by high-resolution melting (HRM) analysis to develop a single closed-tube screening panel to screen for Lynch Syndrome. This comprises tests for microsatellite instability (MSI), MLH1 methylation promoter and BRAF mutation.

Methods
For MSI-testing, 5 mononucleotide markers (BAT25, BAT26, BCAT25, MYB, EWSR1) were developed. In addition, primers were designed to interrogate Region C of the MLH1 promoter for methylation (using bisulphite-modified DNA) and to test for mutations in codon 600 of BRAF. Two separate cohorts from Nottingham (n = 99, 46 with MSI, 53 being microsatellite stable (MSS)) and Edinburgh (n=88, 45 MSI, 43 MSS).

Results
All the cases (n=187) were blind tested for MSI and all were correctly characterised by our panel. The MLH1 promoter and BRAF were tested only in the Nottingham cohort. Successful blinded analysis was performed on the MLH1 promoter in 97 cases. All MSS cases showed a pattern of non-methylation whilst 41/44 cases with MSI showed full methylation. The three cases with MSI and a non-methylated pattern had aberrations in MSH2 and MSH6 expression. BRAF mutation was detected in 61% of MSI cases and 11% of MSS cases.

Finally, 12 cases were blind screened by using the whole panel as a single test. Of these, 5 were identified as MSS, 4 as MSI/non-LS and 3 as MSI/possible LS. These results were concordant with the previous data.

Conclusion
We describe the Nottingham Lynch Syndrome Test (N_LyST). This is a quick simple cheap method for screening for Lynch Syndrome.
INTRODUCTION

Colorectal cancer (CRC) is a leading cause of cancer-related mortality (1-3). Most CRCs arise sporadically without any antecedent family history. There are, however, several cancer syndromes in which development of CRC is part of the phenotypic spectrum (1, 2). The most common of these is Lynch Syndrome (also known as Hereditary Non-Polyposis Colorectal Cancer) which is responsible for 2-4% of all CRCs (3, 4). Patients with LS are susceptible to the development of CRCs and to the development of extra-colonic tumours – most notably endometrial, ovarian and small intestine adenocarcinomas (5, 6).

Lynch Syndrome (LS) arises as a consequence of germline mutation in one of four DNA mismatch repair (MMR) genes (i.e. MLH1, PMS2, MSH2, MSH6) (7, 8). Loss of any of the proteins results in loss of MMR function and an increase in the rate of gene mutation. One of the manifestations of this is an increase in insertion-deletion (indel) mutations especially at DNA microsatellites – known as microsatellite instability ( MSI) (8, 9). Tumours arising in LS therefore usually show both loss of expression of at least one of the MMR proteins (i.e. dMMR) and MSI (10, 11). Thus, dMMR and MSI – although they are distinct phenomena, are usually regarded as synonymous. Conversely, proficient mismatch repair (pMMR) is considered synonymous with a microsatellite stable (MSS) phenotype.

Numerous studies have shown that, due to the high risk of multiple cancers and its relatively high prevalence, there is a clinical and economic benefit to be gained by screening CRCs for LS (12-15). Whilst a definitive diagnosis of LS can only be made by demonstration of a germline mutation in an MMR gene, the possibility of LS can be inferred if a tumour is shown to be dMMR or shown to have MSI. However, approximately 10-15% of sporadic CRCs will also show dMMR/MSI due to somatic loss of MMR function (10). Epigenetic silencing of the MLH1 gene is the most common cause of dMMR in sporadic tumours and very rarely occurs in LS (16, 17). Thus sporadic tumours with dMMR/MSI can be distinguished from tumours arising in LS by demonstrating methylation of the MLH1 promoter. Similarly, somatic mutation of BRAF is common in sporadic tumours with MSI but very rarely occurs in tumours arising in LS (17-19).

Guidance from the National Institute of Clinical and Healthcare Excellence (NICE) recommends that all CRCs should be screened for the possibility of LS (12). The pathway suggested involves two steps: firstly, identify cases with dMMR/MSI and
then filter out sporadic cases by testing for *BRAF* mutation and *MLH1* promoter methylation. For the first step, testing for dMMR can be performed by immunohistochemistry (IHC) whilst testing for MSI involves PCR followed by capillary electrophoresis. For the second step, PCR followed by mutation screening or sequencing is required for detection of *BRAF* mutation. Testing for *MLH1* promoter methylation can be performed by PCR on modified DNA followed by sequencing or gel electrophoresis.

This strategy uses multiple tests and requires downstream analysis of the PCR products on different platforms. We believed that testing could be simplified using High Resolution Melting (HRM) analysis. HRM is an exquisitely sensitive method for detecting variations in DNA sequence (20-22). It can be performed at the end of a PCR without needing to transfer PCR products to another tube (i.e. a closed-tube test). We have shown previously that HRM can be used for testing for microsatellite instability (23, 24), for detection of *BRAF* mutation (23-27) and to identify promoter methylation (25). Here we sought to create a single panel test in which a single PCR run followed by HRM can be used to screen for patients at risk of Lynch Syndrome.

**MATERIALS AND METHODS**

**Cell lines**

CRC cell lines were kindly donated by Prof Ian Tomlinson. The cell lines DLD1, HCT116, RKO, LoVo, and LS1034 have previously been shown to have MSI whilst the cell lines SW480, SW620, HUTU80, SW837 have been shown to be microsatellite stable (MSS) (28). DNA was extracted from cell lines using the Qiagen DNeasy kit (Qiagen, UK) as per manufacturer’s instructions and adjusted to a concentration of 20 ng/µl. Identity of the cell lines was confirmed by mutation profiling as previously described (25).

Two diploid cell lines were chosen for spiking experiments in order to perform limit of detection experiment. DNA extracted from HCT116 (an MSI cell line) was spiked into DNA extracted from SW837 (MSS), to produce mixtures of DNA containing various proportion of HCT116 of ≈50%; ≈25%; ≈12.5%; ≈6%; ≈3% and ≈1.5%.

**Primary colorectal cancers**

*The Nottingham cohort*
Ninety nine cases of CRC, which had previously been tested by immunohistochemistry for expression of MMR proteins, were retrieved from the archives of Nottingham University Hospital (NUH) Pathology Department. Of these, 46 cases which were dMMR (and by inference had MSI). The remaining 53 cases were pMMR (and by inference were MSS). Access to tissues and ethics approval were granted by Nottingham Health Sciences Biobank which has approval as an IRB from North West - Greater Manchester Central Research Ethics CommitteeREC reference: 15/NW/0685.

The Edinburgh cohort
Eighty eight cases of CRC were retrieved from the archives of the Royal Infirmary of Edinburgh, Pathology Department which had previously been tested for expression of MMR proteins or MSI. Of these, 45 cases were dMMR/MSI and 43 cases were pMMR/MSS. Access to anonymised use of tissues was granted by Tissue Governance NHS Lothian under ethics approval number SR783.

DNA extraction for formalin-fixed tissue
DNA was also extracted from formalin-fixed paraffin-embedded (FFPE) tumour samples. One or two 20 µm thick sections (depending on tissue surface area) were cut from each block. DNA was extracted using the QIAamp DNA FFPE tissue kit (Qiagen Ltd, UK) following the manufacturer’s protocol. All DNA samples were adjusted to a concentration of 20 ng/µl.

Validation of immunohistochemistry as a marker of MMR deficiency
The Nottingham cohort had been tested for expression of the MMR proteins by immunohistostaining (IHC). In order to confirm that the interpretation of the IHC was a correct reflection of the MMR function, a group of 33 cases (15 MSI/18 MSS) were tested by PCR followed by capillary electrophoresis (CE). PCR and CE testing was performed by the Molecular Genetics Laboratory at Nottingham University Hospitals NHS trust using the Promega MSI System version 1.2 in accordance with manufacturer’s instructions. Five mononucleotide markers for MSI testing (BAT-25, BAT-26, NR-21, NR-24, MONO-27) (29) and two pentanucleotide markers (Penta-D, Penta-E) for sample identity checking, were amplified using fluorescently-labelled primers in a multiplex PCR. Products were analysed by capillary electrophoresis on
an Applied Biosystems 3130xl Genetic Analyzer (Life Technologies) using the kit internal lane standard. Data were analysed using GeneMapper® software. Samples that showed microsatellite instability at ≥2 mononucleotide loci were interpreted as having MSI.

**Identification of novel markers for MSI testing**

A total of 11 different mononucleotide repeat microsatellite loci with potential utility as sensitive markers of MSI were tested. Of these, two markers (BAT 25 and BAT 26) are established MSI markers although the primers for these loci were redesigned (30-33). Two markers (BCAT 25 and TYMS 26) were identified via bioinformatics analyses by our group (we have previously described BCAT 25 (23, 24) as a useful marker), two markers (EWSR1 and MONO-27) were identified from published research articles (34-36) and five markers (MYB, ANGEL2, TP53 (BAT34CA), FBXO46, and TCF4) were identified in the SelTar database (37) ([www.seltarbase.org](http://www.seltarbase.org)). The individual markers were chosen if (i) the mononucleotide repeat size is >10 bases and (ii) if the published rate of mutation of the marker in MSI tumours (CRC and gastric cancer) is >80%. Primers were designed using a combination of online design tools: MFeprimers ([http://mfeprimers.igenetech.com](http://mfeprimers.igenetech.com)), UCSC in silico PCR ([http://rohsdb.cmb.usc.edu/GBshape/cgi-bin/hgPcr.](http://rohsdb.cmb.usc.edu/GBshape/cgi-bin/hgPcr.)) and Reverse Complement ([www.bioinformatics.org](http://www.bioinformatics.org)). The optimum annealing temperatures of the primer pairs was ascertained as previously described (38). Supplementary Table 1 lists all the mononucleotide repeat microsatellite markers, their genomic locations, amplicon sizes, the lengths of the mononucleotide repeats and the ranges of optimum annealing temperature. A range of metrics were used in order to define the best primers including reproducibility, PCR efficiency and range of functioning annealing temperature.

**Testing for MSI using HRM analysis**

In order to test for MSI using HRM, PCR was carried out each sample on the ABI 7500 FAST Real-Time PCR System (Applied Biosystems). Each reaction was carried out in a final volume of 10µl and contained 5µl of 2x Hot Shot Diamond PCR master mix, 0.5µl of 20x (25µM) EvaGreen dye, each primer final concentration was at 0.25µM and 20ng DNA template. The PCR was performed using a 3-step procedure: 1 cycle of (95°C/5 minutes), 45 cycles of [(95°C/10 seconds)X1; (55°C/30
seconds) X 1; (72°C/30) X 1] and 1 cycle of (72°C/2 minutes). HRM was performed in-
tube immediately after PCR and consisted of heating to 95°C for 15 seconds, rapid
cooling to 60°C and maintenance at 60°C for 1 minute. This was followed by slow
ramping up at 0.03 degrees/s to 95 °C during which fluorescent data were captured.
The melting data were analysed following normalisation but without temperature
shifting using the ABI HRM software v2.0. Samples were regarded as MSI if ≥2
markers (40%) showed instability; otherwise, they were regarded as microsatellite
stable (MSS) tumours.
The limit of detection for MSI by both CE and HRM was tested using spiked DNA
samples (as described above).

**Novel primers for BRAF testing**

We have previously designed primers for screening for BRAF mutation using the
nested QMC-PCR protocol (26, 27). For the purposes of this protocol, which requires
a single stage PCR, novel primers were designed specifically for detection of
mutation at codon 600.

**Testing for methylation of the MLH1 promoter**

*Primer design*

Bisulphite modification of DNA causes a conversion of non-methylated cytosine
residues to uracil whilst the methyl group of the methylated cytosines protects
against this change (and cytosines are preserved). Following PCR on bisulphite
modified DNA, the methylated cytosines remain whilst non-methylated cytosines are
converted to thymine residues. The sequence of methylated / non-methylated DNA
is therefore different and can be discriminated by HRM.

The promoter of *MLH1* contains 4 CpG rich regions (labelled A – D) which are the
targets of epigenetic modification. It is generally considered that hypermethylation of
the CpG island in Region C is related to *MLH1* silencing (39). Furthermore, it is
reported that Region C exists in a dichotomous state i.e. all CpG residues being
either methylated or non-methylated (39, 40) without a state of partial methylation.
However, the exact location of Region C is not well defined and the number of
reported CpG residues varies between 5 and 8 (39, 40). Using the publicly available
data, we identified a part of Region C (located −46 to −111 from the transcription
start site; NCBI sequence ID: NC_018914.2) which would contain all 8 of the
reported methylated CpG residues (Supplementary Figure 1). Primers were designed to interrogate the whole CpG island of Region C using the exactly the same cycling and HRM parameters as for the MSI markers. All tests (both sequencing and HRM) for MLH1 Region C promoter methylation were performed on bisulphite modified DNA.

*Bisulphite conversion of DNA*

In order to test for methylation of the MLH1 promoter, it was necessary to modify the DNA. Bisulfite conversion of 400 ng of genomic DNA from each sample was carried out using the EZ-DNA Methylation-Lightning™ Kit (Zymo Research, USA), according to the manufacturer's protocol. Optimisation of the methylation detection HRM-PCR assay was carried out using completely methylated or non-methylated human control DNA (Qiagen Ltd, UK).

*Sequencing of Region C of the MLH1 promoter*

In order to confirm the dichotomous methylation state of Region C, twenty CRCs (10 pMMR, 10 dMMR) were selected from the Nottingham cohort for Sanger sequencing of modified DNA. PCR prior to sequencing was performed using the reverse primer as described above. The forward primer however was modified to include a “squirrel” tail to allow sequencing of short fragments as previously described (38). PCR products were purified using the QIAquick kit (Qiagen) and the products sent to the DNA sequencing facilities (School of Life Sciences, University of Nottingham) and sequenced using Applied Biosystems BigDye Terminator v3.1 Cycle Sequencing Kit and 3130xl ABI PRISM Genetic Analyzer (Data collection software v3.0, Sequence analysis software v5.2). The chromatograms were interpreted using Finch TV 1.4.0 free software from www.geospiza.com/finchtv.

*Evaluation of the N_LyST panel*

All the biomarkers were tested together as a single panel test. Twelve cases were selected from the Nottingham cohort. This selection contained five cases designated as MSS, four cases designated as MSI with MLH1 deficiency and three cases designated as MSI with deficiency of MSH2/MSH6. They were assigned a new ID and were tested blind.
Statistical analysis
GraphPad Prism software version 5.0 was used for statistical analysis. The Chi squared test was used to test for association between different factors. A value of p<0.05 was taken as being statistically significant.

RESULTS
Validation of IHC as a marker of mismatch repair function
The Nottingham cohort had been selected using IHC expression of MMR proteins as a marker of MMR function. To confirm the association between IHC data and the presence or MSI, 33 cases from this cohort were tested by CE for the presence of MSI. Of these, 15 had been designated dMMR and 18 were pMMR. There was 100% concordance between the IHC analysis and MSI test results.

Utility of HRM for detection of MSI
From 11 different potential microsatellite loci, a panel of 5 markers comprising BAT25, BAT26, BCAT25, MYB and EWSR1 was chosen as the one showing the best performance. Our panel was compared with the commercial CE panel for their limit of detection for MSI calling. Using spiked samples containing varying proportions of DNA from MSI/MSS cell lines, the CE method and HRM were comparable with a limit of detection ≈6.25% (Figure 1).
Our panel was used to test the Nottingham cohort of 99 cases of CRC (46 dMMR and 53 pMMR) and both observers correctly called every case. The Edinburgh cohort of 88 (45 dMMR/MSI and 43 pMMR/MSS) were tested separately. The HRM data were analysed by the same two observers and one observer correctly called all cases whilst the other observer miscalled 2 case of MSI as MSS. Although we applied the generally used threshold of instability at ≥2 markers (40%) for a call of MSI, most cases usually showed instability at 4-5 markers and only one case, out of the total of 91 cases designated dMMR/MSI, was found to have instability at only 2 markers. Of the cases designated as MSS, 7% (7/96) had instability at 1 marker only whilst the remainder did not show any alteration in the microsatellite markers.

Screening for BRAF mutation
New primers to screen for BRAF codon 600 mutation were designed and optimised to work as a single stage test using the cycling conditions for MSI testing. Primers
were optimised and tested on cell lines with known \textit{BRAF} mutation status (data not shown). All cases in the Nottingham cohort were tested and 28/46 (61\%) of cases designated as MSI showed mutation whilst 6/53 (11\%) of the MSS cases showed mutation. This frequency of mutation is consistent with published data and confirm the significant association of MSI with \textit{BRAF} mutation (chi squared test, \(p<0.0001\)).

\textbf{Analysis of MLH1 promoter methylation}

\textit{Sequencing of Region C}

Twenty cases of CRC form the Nottingham cohort (10 MSI, 10 MSS) were tested for \textit{MLH1} promoter methylation by direct sequencing. Our findings replicated published data with 10/10 case of MSS CRC showing conversion of all 8 of the cytosines at the CpG sites to thymine without any cases suggesting partial methylation (i.e. methylation at some residues but not others). In contrast, 10/10 cases of the MSI CRCs showed retention of the cytosines at the CpG sites (Figure 2). The MSI samples did however show a double signal at the CpG sites i.e. a cytosine and a thymine. Since tumour samples contain both tumour epithelium and stroma, it is expected that the methylated signal comes from the tumour cells whilst the signal from the stroma would be non-methylated.

\textit{HRM analysis of Region C}

HRM was performed following PCR with primers targeted to amplify around the CpG island of Region C of the \textit{MLH1} promoter. Amplification, following bisulphite modification, of both fully methylated and fully non-methylated DNA gave a single peak (Figure 3A). The melting temperature (Tm) of the PCR product from the non-methylated DNA (i.e. the “non-methylated peak”) was lower than that of PCR product from the methylated DNA (the “methylated peak”) reflecting the enrichment the latter with cytosine residues within the methylated sequence.

All cases in the Nottingham cohort were tested for methylation of Region C. Two cases (both designated as MSI) could not be tested due to failed PCR post bisulphite modification of DNA. Of the 97 successfully tested cases, two distinct melting patterns were seen i.e. a single peak low Tm peak (corresponding to the non-methylated peak) and a double peak with both low and high Tm (corresponding to both the non-methylated peak and the methylated peak, Figure 3B). All cases
designated as pMMR/MSS showed only a single non-methylated peak i.e. there was no promoter methylation. We regard this as the “non-methylated pattern”. Of the 44 cases with MSI, 41 showed a double peak indicating both methylated DNA and non-methylated DNA. The double peak was associated with loss of MLH1 expression (Chi squared test p<0.0001) and we regard this as the “methylated pattern”. The double peak is mostly likely due to methylated DNA being present in the tumour epithelium whilst the stromal cells are likely to contain non-methylated DNA. The three remaining MSI cases showed a single non-methylated peak. These cases were deficient in MSH2 and MSH6.

Screening for Lynch Syndrome using N_LyST

In order to test the N_LyST panel, 12 cases were blind-tested in a single PCR run. The outcome of N_LyST is to categorise cases “probable Lynch Syndrome” if they show MSI, have wild-type BRAF and have a non-methylated pattern for Region C of the MLH1 promoter. Any other pattern would be categorised as “not Lynch Syndrome”. All cases were correctly identified by the panel (Table 1, Figure 4).

DISCUSSION

In this paper we have described the Nottingham Lynch Syndrome Test (N_LyST) as a single panel closed-tube test for Lynch Syndrome screening. The cases used to develop this test were selected on the basis of MMR protein expression and, to validate the use of these cohorts for our assay, we firstly confirmed that dMMR based on IHC was very strongly correlated with MSI. N_LyST incorporates the three components of LS screening (i.e. testing for MSI, MLH1 promoter methylation and BRAF mutation) into a single PCR run. Firstly we developed a panel of five microsatellite markers which includes two established markers (BAT25, BAT26) and three novel markers (BCAT25, MYB and EWSR1). When tested in 187 CRCs (from two different institutions), there was near perfect concordance with the IHC/CE designation. Analysis of the HRM data was undertaken by two observers thereby demonstrating that the analysis is easy and reproducible. The HRM method has a similar limit of detection as CE analysis (≈6.25% mutant DNA) but CE analysis can be complicated by stutter bands that can cause difficulty in allele sizing (33, 41).
Next we designed an assay to detect mutations in codon 600 of \textit{BRAF}. Reassuringly the detected mutation frequencies (61\% in MSI tumours, 11\% in MSS tumours) were in the expected range and the association of MSI with \textit{BRAF} mutation (Chi squared test, p<0.0001) was seen.

The third step was the design of an assay to test for methylation of Region C of the \textit{MLH1} promoter. Our sequencing and HRM data confirmed the dichotomous state of Region C i.e.

either non-methylated or fully methylated. The HRM assay clearly discriminated the two states and, when tested on the Nottingham cohort, all dMMR cases with loss of MLH1 expression by IHC had \textit{MLH1} promoter methylation (i.e. the methylated pattern of two peaks) and were therefore sporadic tumours. None of the cases which were pMMR or dMMR due to MSH2/MSH6 loss, had \textit{MLH1} promoter methylation. Finally, all components of N-LyST were put together and tested as a panel. Twelve cases of CRC were blind tested and perfectly categorised as “non-LS” or “probable LS”.

\textbf{N\_LyST} involves a panel of seven PCRs which are performed in a single run using a single cycling program. It could hugely improve workflow in a diagnostic lab since HRM is performed in-tube on completion of the PCR and transfer of PCR products to another platform for further analysis is not required. Since the test involves a panel applied to all tumours, it does mean that some tumours which are MSS will be unnecessarily tested for \textit{BRAF} mutation and \textit{MLH1} promoter methylation. However, the cost of this is more than off-set by savings made on manpower and consumable due to the removal of downstream analyses of PCR products. Furthermore, since it is a closed-tube test, the risk of laboratory contamination with PCR products is eliminated.

Most modern Real-Time PCR machines will have HRM capabilities and expensive specialist equipment is not required for \textbf{N\_LyST}. The ease of the methodology and data interpretation mean the N-LyST could probably be performed in non-specialist diagnostic pathology labs. This becomes pertinent when considering that MSI testing is likely to increase as it provides information which extends beyond Lynch Syndrome testing e.g. MSI can be used to stratify patients in to groups eligible for treatment with 5-Fluorouracil based therapy (42, 43) or immunotherapy (44, 45). Such high throughput analysis will require a rapid and simple test such as \textbf{N\_LyST}.

An important question is whether \textbf{N\_LyST} – since it is a screening test - is relevant in the era of Next Generation Sequencing (NGS). The sheer sequencing power of NGS...
platforms would allow the MMR genes and multiple microsatellites to be sequenced in a single test (46). However, microsatellites regions can be problematic from some NGS platforms and, where there is low tumour epithelium content, great sequencing depth may be required. In addition, MLH1 promoter methylation testing would require Methyl-Seq to be performed. The economic analyses performed as part of the NICE guidelines concluded that it was more cost-effective to screen the tumour samples prior to germline sequencing (12). Since N_LyST can be performed in less time than that required for library preparation and sequencing with NGS, a case for including N_LyST in the testing pathway can be made.

In summary, N_LyST is based on PCR and HRM and uses a panel of 7 markers to test for MSI, MLH1 promoter methylation and BRAF mutation in a single PCR run. It can be performed on most Real-time PCR machines and, as a closed-tube test, it can improve laboratory workflow and reduce turnaround times for testing. It is a robust test which represents a quick, cheap and easy way to screen for Lynch Syndrome.

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<tr>
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</tr>
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FIGURE LEGENDS

Figure 1: Limit of detection of MSI screening by HRM analysis and capillary electrophoresis. Cell line genomic DNA from HCT116 (MSI) was admixed with DNA from SW837 (MSS) to produce differing proportion of MSI/MSS DNA with calculated percentage MSI DNA as indicated. Figure 1A shows the analysis by HRM for each marker. A known MSS DNA sample was used as an internal reference standard. Samples containing 6.25% MSI DNA clearly show different melting pattern as compared to the MSS sample. Figure 1B shows the analysis of the same samples using capillary electrophoresis which depends on fragment size analysis. MSI is indicated by novel alleles of the mononucleotide markers with different sizes compared to the normal sample. MSI could be called in samples containing 6.25% MSI DNA.

Figure 2: Sequencing of Region C of the MLH1 promoter. Region C was identified and Sanger sequencing (following PCR of bisulphite modified DNA) was performed on 10 cases of MSI CRC showing loss of MLH1 expression and MSI and 10 cases of MSS CRCs. Figure 2A shows a chromatogram from an MSI CRC. The sequencing trace shows double peaks indicating C and T residues at all eight CpG sites within this region (arrows). The position in relation to the transcription start site is also indicated. All tested MSI CRCs showed this pattern. Figure 2B is a chromatogram from an MSS CRC and shows a single peak (arrows) indicating a T residue at all eight CpG sites. All tested MSS CRCs showed this pattern and the absence of any double peaks would suggest that there is no partial methylation in these tumours.

Figure 3: HRM analysis of Region C of the MLH1 promoter. In order to define the melting patterns of methylated / non-methylated Region C of the MLH1 promoter, PCR was performed on fully methylated or fully non-methylated control DNA following bisulphite modification. Figure 3A is a derivative plot of the PCR products and shows that each condition (i.e. methylated or non-methylated) had a distinct melting peak. The melting temperature (Tm) of the methylated peak (double arrow) was higher than that of the non-methylated peak (single arrow) reflecting the higher proportion of cytosine residues within the fully methylated samples. Figure 3B shows the melting pattern of tumour samples which are pMMR (single arrow) and dMMR due to loss of MLH1 expression (double arrow). The pMMR tumours gave a single
non-methylated peak. The dMMR tumours gave a double peak representing a methylated peak (from the tumour cells) and non-methylated peak (from the stromal cells). All pMMR tumours tested gave a single peak and all dMMR tumours gave a double peak thereby reinforcing the data that this region does not have a state of partial methylation.

**Figure 4:** *Utility of the N_LyST panel to screen for Lynch Syndrome.* Twelve cases of CRC were randomized and tested using N_LyST. Melting plots are shown for BCAT25 (A), BAT25 (B), BAT26 (C), MYB (D), EWSR1 (E) and BRAF (F). Derivative plots are shown for Region C of MLH1 (G). It can be clearly seen that the melt curves of the tumours with MSI are different from those with MSS. If any marker in a tumour with MSI is stable, it will usually melt in the same way as the markers in the MSS tumours. In D there is one case with MSI (asterisked) which melted differently from the rest of the MSS category but also clustered independently from the other six MSI cases, hence this was called equivocal. F shows that the melt curves of the tumours with wild type BRAF are different from those with mutant BRAF. G is a derivative plot and tumours demonstrate two discrete melting forms: “methylated” comprising two melting peaks which represent methylated DNA (from tumour epithelium) and non-methylated DNA (from tumour stroma) or “non-methylated” comprising one melting peak which characterises a completely non-methylated tumour and stroma cell population.

**Table 1:** *Screening cancers for Lynch Syndrome using the N_LyST panel.*

| Twelve cancers from the Nottingham cohort were screened blind for Lynch Syndrome in a single PCR run (see also Figure 4). The left hand side of the table shows the N_LyST data whilst the original data for these cases are given on the right hand side of the table. The mononucleotide markers define whether a tumour has microsatellite instability (MSI) or whether it is microsatellite stable (MSS). The pattern of MSI, non-methylated MLH1 promoter and wild-type BRAF is indicative of probable Lynch Syndrome whilst all other patterns indicate that Lynch Syndrome is not likely. From this series, three cases (shaded) were correctly identified as probable LS and nine as non-Lynch Syndrome (pLS = probably Lynch Syndrome, U=unstable, S = stable, E = equivocal, F = failed, Non-Meth = non-methylated, Meth = methylated, WT |
= wild type, M = mutant, IHC = immunohistochemistry, dMMR = deficient mismatch repair, pMMR = proficient mismatch repair, MLH1- = no MLH1 expression, MSH2 = no MSH2 expression, MSH6- = no MSH6 expression).

Supplementary data

**Supplementary Figure 1:** Mapping the CpG island of Region C of the MLH1 promoter. Using publicly available data, the CpG island of Region C (containing 8 CpG dinucleotides reported to be invariably completely methylated when MLH1 is silenced by promoter methylation. The 8 CpG sites are underlined with green boxes and are located -46 to -111 upstream of the transcription start site (NCBI sequence ID: NC_018914.2). Primer attachment sites are highlighted in yellow.

**Supplementary Table 1:** Characteristics of genomic markers selected for MSI testing
## Supplementary Table 1: Characteristics of genomic markers selected for MSI testing

<table>
<thead>
<tr>
<th>Marker</th>
<th>Genome location</th>
<th>Gene/position</th>
<th>Source of marker*</th>
<th>Published mutation rates</th>
<th>Pcr product size</th>
<th>Repeat Size</th>
<th>Primer Ta range</th>
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