A homogeneous HLA-B*27 genotyping assay using dried reagent mixtures

Minna Kiviniemi\(^{a,\ast}\), Jorma Ilonen\(^{a,b}\) and Timo Lövgren\(^{c}\)

\(^{a}\)Immunogenetics Laboratory, University of Turku, Tykistökatu 6 A, 4th floor, FIN-20520 Turku, Finland
\(^{b}\)Department of Clinical Microbiology, University of Kuopio and Biocenter Kuopio, FIN-70211, Kuopio, Finland
\(^{c}\)Department of Biotechnology, University of Turku, Tykistökatu 6 A, 6th floor, FIN-20520 Turku, Finland

Abstract. The presence of HLA-B*27 allele with patients suspected with ankylosing spondylitis can be used in the diagnostic process. We have developed an assay for typing for the HLA-B*27 in whole blood dried on sample collection cards using pre-dried reagent wells and homogeneous time-resolved fluorescence based PCR approach. Essentially only the sample needs to be added to the dry ready-to-use reaction well in order to start the homogenous amplification assay. The method was validated with 229 samples also typed with an existing DELFIA-based method and results of both assays were 100% concordant. The dried reagents were shown to be stable at least up to eight weeks at room temperature without any decline in their performance.

Keywords: Dry chemistry, HLA-B*27, homogeneous assay, PCR

1. Introduction

The major application of genotyping for human leukocyte antigens (HLA) is identification of compatible donor recipient pairs for tissue transplantation. Demanding high resolution full-house approaches, with as detailed genotyping of an individual as possible, are commonly used to ensure as complete identity as possible. Besides tissue transplantation definition of certain HLA alleles and allele combinations is used for diagnostic purposes making use of the known disease associations. Typing of only one or few alleles possibly at low level of resolution allows application of methodologies more suitable for handling of large sample numbers and automation.

The association of HLA-B*27 with ankylosing spondylitis (AS) is one of the strongest associations and the allele is also found increased among several types of postinfectious reactive arthritides; around 90% of AS patients carry this allele compared to approximately 9% of controls [1,3,11,17]. Typing for the presence of HLA-B*27 is thus widely used in diagnostic laboratories using a multitude of methods including serological and different sequence based techniques such as PCR with sequence-specific primers [14], PCR-RFLP-analysis [13] and ligation based typing [4].

We have previously developed a homogeneous assay approach for genotyping utilizing asymmetric PCR and lanthanide labelled DNA-probes with LNA bases [8, 9]. Homogeneous assays are separation-free methods were the unreacted label need not be removed from the reaction before measuring the results. The method has been adapted to both extracted DNA samples as well as whole blood samples dried on sample collection cards, the latter being more convenient with the exclusion of a tedious DNA-extraction step [7]. The benefits of this homogeneous approach include reductions in hands-on time as well as whole assay time, reduced contamination risk and, as a concerted consequence of reduced time and material, reduced costs.

The aim of the study was to apply this homogeneous assay principle to an assay for testing the presence of HLA-B*27. We compared the method with an existing HLA-B*27 assay based on DELFIA technology (PerkinElmer Life and Analytical Sciences, Wallac) which we have developed and earlier found to perform...
favourably compared to traditional methods. Also the suitability of the homogeneous HLA-B*27 assay for a dry reagent approach, consisting of drying all the reagents excluding the sample into the reaction vessel, was tested in order to further increase the simplicity of the assay performance since only the sample with water would then need to be added when performing the assay.

2. Materials and methods

2.1. Oligonucleotides

The oligonucleotides used in this study and described in Table 1 were purchased from Thermo- Electron GmbH (Ulm, Germany) except the probes used in the DELFIA-assay which were purchased from PerkinElmer Life and Analytical Sciences Wallac (Turku, Finland). The other lanthanide probes were labelled at Abacus Diagnostica Ltd. (Turku, Finland).

2.2. DNA samples

The EDTA-treated blood was applied to FTA® Classic Card (Whatman International Ltd., Maidstone, UK) sample collection cards and allowed to dry at room temperature. 3 mm discs were cut from the dried cards into the wells of a ThermoFast® 96 PCR plate (ABgene Limited, Epsom, UK). 50 µl of H2O was added into each well and the plate sealed with a Microseal ‘A’ Sealing film (Bio-Rad Laboratories, Hercules, CA, USA). The plate was incubated at 100°C for ten minutes, cooled to 10°C and centrifuged briefly to collect all the liquid to the bottom of the wells. 1 µl samples of the liquid phase containing DNA liberated from the blood discs were used as template in the PCRs. The concentration of the released DNA was not measured as the assay was optimized to the amount available from blood spots and no indication was found that the concentration would be a critical factor.

2.3. The homogeneous HLA-B*27-assay

The PCR components except the DNA polymerase were dried in an Eppendorf tube in vacuum concentrator (Heto-Holten, Denmark). 2 U of DyNAzyme II DNA polymerase (Finzymes, Espoo Finland) were air-dried into each well of a ThermoFast® 96 – PCR-plate (skirted, low profile, black), except for the amplification background wells which did not contain the enzyme. The PCR mix was reconstituted with water and 25 µl of the reconstituted mix per well was added onto the PCR plate containing the dried polymerase. The reconstituted reactions contained 4.0 mM MgCl2, 1x DyNAzyme™ reaction buffer, 0.2 mM dNTPs (Fermentas Inc., Helsingborg, Sweden), 0.4 µM β-actin 5’-primer, 0.08 µM β-actin 3’-primer, 0.1 µM HLA-B*27-specific 5’-primer, 0.2 µM HLA-B*27-specific 3’-primer and 3.0 µg/µl BSA (Sigma-Aldrich Finland Oy, Helsinki, Finland). The concentrations of the probe-quencher-pairs were for the β-actin 3.3 nM (probe) and 6.6 nM (quencher) and for the HLA-B*27 0.33 nM (probe) and 5.0 nM (quencher).

After 1 µl samples of the DNA liberated from the dried blood spots were added, the plates were sealed with Microseal ‘A’ film and a PCR program was run in a DNA Engine Dyad Peltier Thermal Cycler (Bio-Rad Laboratories) consisting of an initial cycle of 4 min at 95°C, 2 min at 64°C, and 2 min at 73°C. This was

<table>
<thead>
<tr>
<th>Oligonucleotide</th>
<th>Sequence from 5’ to 3’ end</th>
<th>Label/position</th>
</tr>
</thead>
<tbody>
<tr>
<td>B*27 5’ primer</td>
<td>GTGGGCTACGTGGACGACACGCT</td>
<td></td>
</tr>
<tr>
<td>B*27 3’ primer</td>
<td>CAGTCTGTGGCGCGTTGCC</td>
<td></td>
</tr>
<tr>
<td>β-actin 5’ primer</td>
<td>TGAAGTGTGACGTGGACATC</td>
<td></td>
</tr>
<tr>
<td>β-actin 3’ primer</td>
<td>CTGATCTCATATTGTGCGG</td>
<td></td>
</tr>
<tr>
<td>B*27-probe</td>
<td>CCATGTGGCATAGAGC 20*Eu/5’</td>
<td></td>
</tr>
<tr>
<td>B*27 quencer</td>
<td>ATCCACGGCdabcyl/3’</td>
<td></td>
</tr>
<tr>
<td>β-actin probe</td>
<td>AGGGTAGCATG</td>
<td>Eu/5’</td>
</tr>
<tr>
<td>β-actin quencher</td>
<td>CATGTACCCT dabcyl/3’</td>
<td></td>
</tr>
<tr>
<td>DELFIA-B*27 5’ primer</td>
<td>GCTACGTGGACGGACGCT</td>
<td></td>
</tr>
<tr>
<td>DELFIA-B*27 3’ primer</td>
<td>CAGTCTGTGGCGCGTTGCC</td>
<td></td>
</tr>
<tr>
<td>DELFIA-β-actin 5’ primer</td>
<td>TGAGTCTGACGTGGACATC</td>
<td></td>
</tr>
<tr>
<td>DELFIA-β-actin 3’ primer</td>
<td>CTGATCTCATATTGTGCGG</td>
<td></td>
</tr>
<tr>
<td>DELFIA-B*27-probe</td>
<td>GCCGTGGCATAGAGC 20*Eu/5’</td>
<td></td>
</tr>
<tr>
<td>DELFIA-β-actin-probe</td>
<td>ATGCCAGGGGTACATG</td>
<td></td>
</tr>
</tbody>
</table>

underlined bases are LNA
followed by 49 cycles of 30 s at 95°C, 40 s at 64°C, ramping of 0.5°C/s to 73°C, and 40 s at 73°C. After a final extension step of 4 min at 73°C, the reactions were slowly cooled to room temperature at an approximate rate of 0.02°C/s. All reactions for altogether 229 samples were done in duplicate.

After thermal cycling and cooling, europium and terbium signals were measured at room temperature with a Victor 1420 Multilabel counter (PerkinElmer Life and Analytical Sciences Wallac). Default parameters of the instrument were used for both europium and terbium measurements (for europium excitation and emission wavelengths 340 nm and 615 nm, respectively, delay 0.4 ms, window time 0.4 ms, and measurement cycle time 1 ms; and for terbium excitation and emission wavelengths 340 nm and 545 nm, respectively, delay 0.5 ms, window time 1.4 ms, and measurement cycle time 2 ms).

To analyze the results, signal-to-background ratios were calculated for both probes by dividing the signal measured from the sample by the signal obtained from a background reaction. For the amplification control the background signal was obtained from a reaction that did not contain the enzyme; and for the HLA-B*27-probe a reaction with a negative control sample was used as background. To eliminate variations caused by differences in amplification efficiency, a relative HLA-B*27 ratio was calculated for each sample by dividing the signal-to-background ratio of the HLA-B*27-probe by that of the amplification control probe.

2.4. The DELFIA HLA-B*27-assay

The 229 samples were also genotyped with the DELFIA-based method published previously [20]. The DELFIA method consists of a PCR-step used to amplify allele-specifically the HLA-B*27 if present in sample and a section of the β-actin gene as an amplification control, followed by a hybridisation step where the PCR-products are collected onto a microtitration well and lanthanide labeled probes are used to detect these PCR-products. In short, 25 µl PCR-reactions consisted of 4 mM MgCl₂, 0.8x ABgene PCR-buffer IV (ABgene), 0.4 µM β-actin control primers, 0.1 µM HLA-B*27-specific primers, 1.5 M betaine (Sigma-Aldrich Finland Oy), 0.2 mM dNTP’s and 0.06 U/µl of ABgene Taq-DNA polymerase. 1 µl samples of the extract from the blood spots were added as template per reaction.

The same thermal cycling protocol was used as for the homogeneous assay except for the annealing temperature which was 62°C instead of 64°C and after the final extension step the reactions were rapidly cooled to 10°C omitting the slow cooling.

After the amplification step the DELFIA plate hybridization was performed. The biotinylated PCR-products were collected to a streptavidin-coated microtitration plate (Kaivogen Oy, Turku, Finland) and rendered single-stranded by denaturing with sodium hydroxide solution and removing the complementary strand by washing. A probe labeled with europium recognizing the HLA-B*27 PCR-product and a probe detecting the β-actin product labeled with samarium were added and allowed to hybridize to their target sequences and unbound label was removed by washes. The signals were developed and measured and the results of the DELFIA method were calculated as described for the homogeneous assay.

2.5. The stability of the dried reagents

The stability of the dried reagents was tested with 0.2 M trehalose (Sigma-Aldrich Finland Oy) as a preserving reagent at room temperature for up to six weeks of storage. The reagents consisting of identical components as described in the homogeneous HLA-B*27 assay with the inclusion of 0.2 M trehalose but without BSA were air-dried onto Thermo-Tube strips (AB-0264, ABgene). The DNA-polymerase was air-dried on separate Thermo-Tube strips and sealed into the storage packages together with the reagent strips.

The possibility of including the DNA polymerase in the dried reagent mix was also tested by air-drying the enzyme together with all the other PCR components. All the PCRs were otherwise done as described earlier for the homogeneous HLA-B*27 assay.

3. Results

3.1. The homogeneous HLA-B*27-assay compared to the DELFIA HLA-B*27-assay as a reference method

Altogether 229 consecutive patient samples tested for the presence of HLA-B*27 using DELFIA method were used to evaluate the performance of the homogeneous assay. Samples were tested in 6 different runs of DELFIA assay with an initial success rate of 84% and the 36 samples that failed to amplify in the first attempt were successfully typed later. The results of the homogeneous assay are shown in Fig. 1. The samples were analyzed in 5 separate runs with an initial success
rate of 96% and the relative signal ratios calculated as described in Material and Methods for all samples are shown in Fig. 1. The nine samples that failed to reach the limit set for acceptable amplification in the first attempt (not included in the figure) were reanalysed successfully.

All samples were successfully genotyped and also were 100% concordant when the results of both assay methods were compared.

3.2. The stability of the dried reagents

The possibility of including the polymerase in the PCR reagent mix was explored and the results are shown in Fig. 2. The relative HLA-B*27-ratios of those reactions where the enzyme was dried separately are clearly divided into HLA-B*27-positive and – negative samples but when the enzyme was dried together with the rest of the PCR components the positive and negative samples can not be distinguished.

The stability of the dried reagents was initially studied using BSA as a stabilizing agent; but BSA did not prove to be effective enough preserving agent when stored at room temperature, at +4°C or at −20°C for several weeks (results not shown). Trehalose was subsequently tested to be used instead of BSA.

The dry PCR reagents with trehalose were stored at room temperature up to eight weeks. The results of the
Fig. 3. The results of the dry chemistry reagent stability tests. Panel A shows the signal-to-background ratio of the europium-labeled amplification control probe and panel B the signal-to-background ratio of the HLA-B*27 probe labeled with terbium at six time points from one day to eight weeks of storage. The cut-off values for the amplification control probe and for the HLA-B*27-probe were set at 1.3 and at 1.5, respectively and are marked with solid horizontal lines in the figure. The B*27-positive samples are marked with filled circles and the B*27-negative samples with open circles.

stability test are shown in Fig. 3 with four HLA-B*27-positive and three HLA-B*27-negative samples included in each time point. All samples were successfully typed at each time point with clearly differing positive and negative HLA-B*27-signals as shown by the panel B in Fig. 3. Although at four weeks one HLA-B*27-positive sample was slightly below the limit set for the control probe signal-to-background ratio as acceptable amplification as shown in panel A in Fig. 3, but despite this it showed an unambiguously positive HLA-B*27 result. The signal-to-noise ratios of both probes do not show any clear declining trend during the eight weeks and even the raw signals obtained from the reactions (data not shown) remain essentially on the same level at all time points.

4. Discussion

We present here a homogeneous method developed for the screening of blood samples for the presence of HLA-B*27 that is simple, robust and fast and also suited for large-scale screening and compares favorably with the previously used DELFIA based assay. In addition the homogeneous method was proven to be adaptable to a dry reagent approach which would fur-
ther simplify the assay and reduce required hands-on
time.

The primers used in the assay have been previously
published [14] and are used also in the DELFIA ass-
say [20]. The primers amplify all HLA-B*27-alleles
shown to be associated with AS; only few rare alleles
are not amplified. A minor modification was done to
the HLA-B*27-specific 5'-primer by adding four bases
to the 5'-end of the primer to compensate for the dimin-
ished concentration of this primer used in the asym-
metric PCR when compared to the HLA-B*27-specific
3'-primer concentration, since the \( T_m \) of the primers
depends on the concentration as well as the length of
the primer [8,15]. The inclusion of LNA-bases in the
lanthanide-labeled probes was shown by our previous
work to improve significantly the performance of the
probes, when compared to regular, only-DNA contain-
ing oligonucleotides, in the developed assay format [8]
and although the specificity in this assay relies mainly
on the HLA-B*27-primers, given that the HLA-B*27
probe used in the homogeneous assay has a sequence
which can be found also in other HLA-B-alleles, such
as HLA-B*07, *08 and *39 among others, we chose to
take advantage of the strong affinity of LNA-bases [2,
12] in this assay as well to allow the use of short probes.

The developed method was optimized to ensure the
slightly preferential amplification of the HLA-B*27
over the \( \beta \)-actin amplification control to guarantee that
no false negative results would be produced. In some
cases even though the signal-to-background ratios of
the control probe did not reach the set cut-off value,
indicating insufficient amplification, the HLA-B*27-
probe showed a clear positive result (data not shown),
indicating that the chosen conditions truly favor the
amplification of the HLA-B*27 over the amplification
control.

Compared with the DELFIA method the interassay
variation was smaller with the homogeneous assay en-
abling easy comparison of separate assay runs, although
all the results were also easily interpreted within each
of the DELFIA assays. Also the initial success rate of
the developed homogeneous method is superior to that
of the DELFIA method demonstrating a more robust
approach.

A dry reagent approach would simplify even further
the developed method since only the pipetting of the
samples and possibly adding of water would be re-
quired. With this in mind the stability of the PCR com-
ponents in dry format was tested with an air-drying pro-
cess that requires no specialized equipment. The time
required for preparing the dry reagents consists of ap-
proximately 30 minutes of hands-on time for preparing
the reagents and dispensing them into the reaction ves-
sels and of overnight drying after which the reagents are
ready to use or can be packaged for storage. With BSA
as a stabilizing agent the stability of the dried reagents
was not sufficient and therefore we tested trehalose that
has been shown to protect dried reagents [10] as well as
act as a PCR enhancer [16,18]. The reagents air-dried
together with 0.2 M trehalose with separately dried
DNA-polymerase were shown to be stable at least up to
eight weeks of storage at room temperature. Since the
signal-to-noise ratios of either the HLA-B*27-probe or
the amplification control probe did not show any clear
decline during this period it could be assumed that even
longer storage time could be used, and these tests are
continuing in our laboratory. Storing the reagents in
lower temperatures, such as at +4°C or even at −20°C,
would likely result in even longer self-life than expect-
ed for storage at room temperature [5,6].

When the possibility of including the polymerase
with the other reaction components was tested, the re-
sults showed clearly that the enzyme could not be dried
together with the other PCR components. However,
the enzyme might be dried in the same reaction vessel
with the other components provided that the enzyme
remains physically separate from the other components
during all of the time of storage [19]. For example,
the enzyme could be dried to a different position in the
PCR vessel while the rest of the mix was on the bottom,
but this would require more sophisticated instrumen-
tation than were available for this study. When this
is accomplished only the sample needs to be added to
the reaction well in order to perform the homogeneous
amplification assay and the assay can be run in a few
hours or less depending on the speed of the thermal
cycler available.

The homogeneous approach also proved to be some-
what cheaper than the DELFIA format when the reagent
and material costs were calculated. Much of the di-
minished costs of the homogeneous assay are due to
the absent hybridization step since the probes used to
recognize the PCR-products are already included in the
PCR mix and this is true also when compared to other
methods used to type for the presence of HLA-
B*27 requiring a separate detection step, such as plate
hybridization or gel electrophoresis. Also the loss of
reagents is smaller if pre-dried reagents are used, since
usually the mixtures used in PCR and hybridization are
prepared with some excess to facilitate pipetting. More
importantly, the time saved in hands-on time with the
homogeneous method reduces the costs of the assay
even more significantly, since labor is in many occasions the most expensive element. Since the separate hybridization step, requiring the greatest part of the hands-on work for example in the DELFIA method, is omitted from the homogeneous method the whole assay can be run in approximately 4 hours with 1 hour or less of hands-on time compared to the minimum of 8 hours required by the DELFIA method with more than 3 hours of hands-on time.

The flexibility of the developed homogeneous HLA-B*27 assay allows either the typing of only a few samples at a time or it can be applied to the screening of hundreds of samples, according to needs of the user, without any modifications to the assay protocol. With the minimized assay and hands-on time, especially when using the pre-dried reagents, and the reduced total costs, the presented method compares favorably with the existing assays in the typing of samples for the presence of HLA-B*27.

References

Submit your manuscripts at http://www.hindawi.com