Detection of adventitious viruses from biologicals using a broad spectrum Microbial Detection Array


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Crystal Jaing\textsuperscript{1*}, Shea Gardner\textsuperscript{2}, Kevin McLoughlin\textsuperscript{2}, James B. Thissen\textsuperscript{1} and Tom Slezak\textsuperscript{2}

\textsuperscript{1}Physical & Life Sciences Directorate, Lawrence Livermore National Laboratory, Livermore, CA 94550
\textsuperscript{2}Computations Directorate, Lawrence Livermore National Laboratory, Livermore, CA 94550

\textsuperscript{*}Corresponding author: Crystal Jaing
Mail address: Lawrence Livermore National Laboratory
Mail Code L-452
7000 East Ave.
Livermore, CA 94550
Phone: (925) 424-6574
Fax: (925) 422-2282
Email: jaing2@llnl.gov

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ABSTRACT:

We designed the Lawrence Livermore Microbial Detection Array (LLMDA), which contains 388,000 DNA probes. This array can detect any sequenced viruses or bacteria within 24 hours. In addition, the oligonucleotide probes were selected to enable detection of novel, divergent species with homology to sequenced organisms. We recently used this array to identify an adventitious virus from a vaccine product. We have also used this array to detect viral and bacterial infections from various human clinical samples. Broad spectrum microbial detection microarrays are efficient and cost-effective tools to rapidly screen cell bank samples, raw materials, vaccine samples and clinical samples to ensure drug, food and health safety in the United States and worldwide.

KEYWORD:

Microarray, adventitious, virus, safety, vaccine, clinical
Introduction

Rapid detection and characterization of bacterial and viral pathogens is important for biodefense and public health. The currently available detection techniques require multiple assays to screen for different markers. A more comprehensive and higher throughput method will greatly improve the efficiency and speed of sample analysis. At Lawrence Livermore National Laboratory (LLNL), we have developed several types of high density microarrays to characterize a broad range of microbial sample constituents, with typical processing times under 24 hours. The Lawrence Livermore Microbial Detection Array (LLMDA) contains 388,000 DNA probes designed to detect any of the currently sequenced viruses or bacteria (1). The LLMDA probes are selected to enable detection of novel, divergent species or strains with sufficient DNA homology to sequenced organisms. We recently used this array to identify contaminating porcine circovirus DNA in a vaccine used worldwide in infants to prevent rotavirus infection (2). We have also used this array to detect viral infections from various human clinical samples (3). We developed a maximum likelihood analysis method that enables us to identify multiple viral and bacterial pathogens from the same sample (1). The array is highly sensitive and able to achieve low-copy bacterial and viral genome detection when coupled with whole genome amplification. Our microarrays are efficient and cost-effective tools to rapidly characterize emerging, unknown and engineered pathogens. The arrays can potentially be used for public health, product and food safety and global disease surveillance and can be further customized for specialized need of these missions.
**Current pathogen detection techniques**

Current detection systems, such as polymerase chain reaction (PCR) technologies, focus on small, prioritized sets of high-risk biological pathogens. PCR analysis is relatively inexpensive and fast. It also has a high sensitivity for known organisms, but it can process no more than about 50 DNA signatures at one time. The likelihood of discovering unknown agents or new species is low with PCR analysis. At the other end of the cost spectrum, high-throughput sequencing provides the most comprehensive information about biological pathogens, both known and unknown. The process, however, is costly and can take several days to produce results.

Microarrays, however, can identify a broad range of organisms, including pathogens on a priority screening list, sequenced bacteria or viruses that might not be anticipated, or even emerging pathogens containing DNA sequences previously identified in other pathogens. Processing samples on an array such as the LLMDA is less expensive and faster than sequencing, and more inclusive than PCR analysis. Figure 1 is a comparison of PCR, microarray and sequencing in speed, cost and resolution.

Several broad spectrum pathogen microarrays have been developed to date. Previously developed microarrays either use small subunit RNA probes for low-resolution identification of bacteria (4), have probes for viruses only (5), or use a combination of these approaches (6). The LLMDA, by contrast, has several dozen probes for each of the thousands of bacteria and viruses sequenced to date, so it can simultaneously examine multiple genomic regions from each organism. Thus, the LLMDA is the only array to date that can provide high-resolution simultaneous characterization of both bacteria and viruses present in a complex sample.
LLMDA probe design for broad spectrum detection of viruses and bacteria

The original LLMDA design covers all bacteria and viruses with sequenced genomes available in spring of 2007, including 38,000 virus sequences representing 2200 species, and 3500 bacterial sequences representing 900 species (1). Updated versions of the LLMDA have been developed to include more recently sequenced genomes. In designing probes for our array, we sought to balance the goals of conservation and uniqueness, prioritizing oligo sequences that were conserved, to the extent possible, within the family of the targeted organism, and unique relative to other families and kingdoms. We designed arrays with larger numbers of probes per sequence (50 or more for viruses, 15 or more for bacteria) than previous arrays having only 2-10 probes per target (5,6). The large number of probes per target was expected to improve sensitivity, an important consideration given possible amplification bias in the random PCR sample preparation protocol, which could result in non-amplification of genome regions targeted by some probes. Probes were selected to avoid sequences with high levels of similarity to human, bacterial and viral sequences not in the target family. Favoring more conserved probes within a family enabled us to minimize the total number of probes needed to cover all existing genomes with a high probe density per target, enhancing the capability to identify the species of known organisms and providing the possibility to detect unsequenced or emerging organisms. Strain or subtype identification was not a goal of probe design for this array. Nevertheless, our ability to combine information from multiple probes in our analysis made it possible to discriminate between strains of many organisms for which multiple genomes are available.
LLMDA processing and analysis

LLMDA processing begins by purifying DNA or RNA from a clinical, environmental or cell culture sample. The purified DNA or RNA is labeled with a fluorescent dye and then hybridized overnight to the microarray, in an incubator heated to 42°C. After we wash off unbound DNA, we scan the array with a laser to excite the labeled DNA targets that are hybridized to matching DNA probes on the array, and measure the fluorescence signal.

To find a combination of organisms whose presence in the sample best explains the observed data, we developed the composite likelihood maximization algorithm (CLiMax), as described in (1). The likelihood maximization algorithm is an iterative process, in which we scan repeatedly through a database of all sequenced microbial genomes. In the first iteration, we look for the target genome that explains the largest portion of the observed detected probe signals, while minimizing the number of probes expected to bind that target that did not have detectable signals. In each subsequent iteration, we choose the target that explains the largest part of the signal not already explained by the first target, while again minimizing the number of expected negative probes. The process continues until a maximal portion of the observed probe signals are explained, or for a specified maximum number of iterations.

The CLiMax algorithm is implemented as part of an automated software pipeline, which incorporates web-based data entry, sample tracking and analysis. The microarray results are shown in a graphical format showing a list of predicted targets organized by viral or bacterial family, together with the log-likelihood scores for each target sequence (Figure 2).
**Identification of adventitious agents from vaccines using the LLMDA**

In early 2010, our team used the LLMDA to evaluate seven live attenuated virus vaccines including Varivax, MMR-II, Attenuvax, Meruvax, Rotateq, Rotarix, and YF-Vax (2). We combined massively parallel 454 sequencing with LLMDA microarray analysis to evaluate any potential adventitious agents in these vaccines; additionally, an oral polio vaccine (OPV) was tested by 454 sequencing only. The expected attenuated viruses were detected in all vaccines tested. The non-vaccine viruses identified from 454 sequencing and LLMDA analysis are listed in Table 1, together with the cell lines used to produce each vaccine. Both 454 sequencing and LLMDA analysis identified the presence of porcine circovirus (PCV) DNA in the Rotarix rotavirus vaccine. The 454 analysis indicated further that a large fraction of the sequence reads from the Rotarix vaccine (41.6%) consisted of PCV1 sequences. Because serum-free media are used in the manufacture of Rotarix, a possible source for PCV1 DNA is porcine pancreas-derived trypsin used for passaging Vero cells. After introduction of PCV1, a chronic infection may have become established in Vero cells (7). Alternatively, since the porcine-derived trypsin is typically irradiated before use to inactivate adventitious viruses, the PCV1 DNA may simply reflect carryover of noninfectious, lethally irradiated PCV1 (2).

In vaccines grown in chicken embryo fibroblast (CEF) cells, signals to avian leukosis viruses (ALV) and avian endogenous retrovirus (AEV) were detected by LLMDA. The detection of ALV and AEV in live attenuated vaccines derived from CEF has been reported previously and extensive investigations indicated these viruses were non infectious to humans (8,9).

The detection of SRV DNA in the Rotateq vaccine by 454 sequencing reflects the presence of endogenous SRV DNA in the germ line of African green monkeys, from which the Vero cell line used to produce the vaccine is derived. SRV is thought to be a highly prevalent
infection of old-world monkeys and to have endogenized at multiple copies in the genomes of many nonhuman primate species (2,10,11). The removal of SRV nucleic acid from the Rotateq vaccine by DNase indicated that it was present as naked DNA released from Vero cells (2). The detection of a retroviral inactivating mutation in the Rotateq SRV DNA pol gene also indicated that this endogenous retrovirus is likely defective (2). LLMDA detected baboon endogenous retrovirus (BEV) sequences from Rotateq. The origin of the BEV signal for Rotateq is probably related to the African green monkey-derived Vero cell used in its manufacture and cross-hybridization of its endogenous retroviruses to the baboon endogenous retrovirus probes. The detection of human endogenous retrovirus K (HEV) in Varivax, MMR-II, and Meruvax was the expected consequence of their manufacture using human cell lines (Table 1).

Clinical diagnostic applications of LLMDA

LLMDA can be used for any applications where the rapid determination of major bacterial and viral components is desired. Recently, we have established a diagnostic platform for unbiased random amplification and microarray identification of viral pathogens in clinical samples (3). We show that Phi29 polymerase-amplification of a diverse set of clinical samples generates enough viral material for successful identification by the LLMDA, demonstrating the potential of the microarray technique for broad-spectrum pathogen detection in human samples. Our studies showed that LLMDA detects both DNA and RNA viruses as well as bacteria and plasmids, present in the same sample, and in some cases can differentiate different subtypes.
Potential of microarray technologies for QC of biological material

High-throughput sequencing and microarrays provide effective means to interrogate current and future vaccines for genetic variants of the targeted viruses and the presence of adventitious viruses. The wide range of sequences detected by these methods, including endogenous retroviruses, bacterial DNA, adventitious viruses such as PCV1, and other nucleic acids of undetermined taxonomic origin, is an expected result given closer scrutiny to the nucleic acids present in vaccines, and not necessarily a reflection of unsafe products.

Advantages and limitations of microarrays

Compared to PCR assays and high-throughput sequencing, microarrays occupy an intermediate range of cost, processing time, and sensitivity for detection of microbial DNA in complex samples. While microarrays are not as sensitive or inexpensive as PCR, they can query hundreds of thousands to several million regions of DNA in parallel (depending on platform and format), compared to at most a few tens of regions in the largest multiplexed PCR assays. Given that approximately 1000 species of human viral and bacterial pathogens have been sequenced at present, PCR assays are too limited for broad-spectrum microbial analysis. However, PCR will continue to be, for the foreseeable future, the most sensitive discriminator of whether a known pathogen is present. Similarly, sequencing will remain the best way to determine the exact nucleotide composition for a novel or mutated pathogen. Nevertheless, we expect that microarrays will evolve during this decade to become the most cost-effective means for quality control of biological products and identifying known pathogens present in clinical samples.

As with any technology based on nucleic acid detection, the capabilities of the LLMDA are limited by the genome sequence information available at the time of design. Many bacterial
and viral species are known but have not yet been sequenced. The LLMDA cannot identify bacteria from families that have absolutely no genomes sequenced, nor can our probe design strategy avoid selecting probes that may cross-hybridize to unsequenced organisms. We are currently seeking funding to update the LLMDA design to take advantage of recent genomic sequence data. Random amplification used with our array may yield lower sensitivity than arrays targeting a limited number of specific regions, for which specific PCR can be used. On the other hand, while specific PCR increases sensitivity for a limited number of targets, it also limits the breadth of targets that can be detected.

We note that United States government agencies have yet to define formal validation procedures for high information content assays such as microarrays and genomic sequencing. This regulatory barrier may delay the widespread use of high-throughput techniques to identify pathogens in human clinical samples, food, vaccines and other products. However, food and pharmaceutical companies and diagnostic laboratories in the US and Europe have shown strong interest in applying these technologies, and we are confident that, with their backing, the attendant regulatory hurdles can be overcome.

Summary

Microarrays provide a cost-effective way to determine the presence of pathogens with known sequence at strain resolution in complex samples, without the larger cost and time required for full genomic sequencing. The LLMDA combines innovative large-scale bioinformatics with modern oligonucleotide array technology to provide the first high-resolution pan-microbial microarray that accurately determines both the bacterial and viral content of unknown clinical or environmental samples. We have demonstrated how this technology can
improve medical product safety (2) and identify viral infections in human samples (3). Future uses of the LLMDA, once permitted by regulatory agencies, will create further cost savings and improvements in research, medical diagnosis, food and product safety applications.

Acknowledgement

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Conflict of interest declaration

A patent application has been filed for the LLMDA design and analysis approach.

References


Table 1. Nonvaccine viruses identified by 454 sequencing and LLMDA

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ALV: avian leukemia virus
AEV: avian endogenous retrovirus
HEV: human endogenous retrovirus
SRV: simian endogenous retrovirus
BRV: baboon endogenous retrovirus
PCV: porcine circovirus
Figure legends

Figure 1. Comparison of PCR, microarray and high-throughput sequencing technologies. PCR, microarray and sequencing are three of the most used DNA detection technologies. PCR is fast and cheap but limited to a few known agents. DNA sequencing has the highest resolution, but the cost is high and it takes a minimum of three days to get results. Microarrays fill some gaps between PCR and sequencing. It has medium cost and results can be obtained within 24 hours. Microarrays can analyze known pathogens and discovery emerging and unknown pathogens.

Figure 2. An example of the LLMDA data from a human clinical sample. The LLMDA was analyzed using the CliMax algorithm developed at LLNL. The results are shown in a list of predicted targets organized by viral or bacterial family, together with the log-likelihood scores for each target sequence.