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Immuno-Oncology Is Making Pharma Step Up Its Diagnostics Game

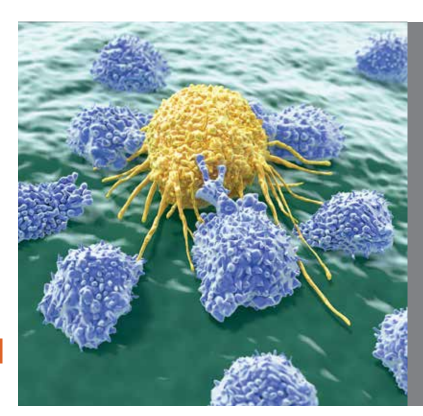
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BY ALEX VADAS

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Immuno-Oncology Is Making Pharma Step Up Its Diagnostics Game



Immuno-oncology has significant diagnostic needs including identifying patients most likely to respond to therapies. Current diagnostic approaches often fall short, due in large part to the complexity of the biology driving therapy responsiveness. As such, the biopharma industry is exploring many emerging approaches covering tumor, immune and even microbiomerelated pathways and biomarkers.

BY ALEX VADAS

Our understanding of the biology driving IO therapy responsiveness is being accelerated by next-generation sequencing profiling of solid tumors linked to clinical and outcomes data, with tumor mutational burden being the prime example.

The acceleration of scientific breakthroughs in IO biology, pathways and biomarkers may simply outpace traditional pharma development.

So what? The pharma industry will need to grapple with what type of biomarker strategy to pursue in IO. Standardization will be key. Pharma must also work with industry partners to implement tests that fit better into therapeutic decision-making windows and health care economics. mmuno-oncology represents a paradigm shift in cancer treatment, with the first wave of PD-1/PD-L1 checkpoint inhibitors such as *Opdivo* (nivolumab, from **Bristol-Myers Squibb Co.**) and *Keytruda* (**Merck & Co. Inc.**'s pembrolizumab) demonstrating cure-like performance in selected metastatic tumors such as non-small cell lung cancer (NSCLC) and melanoma.

This efficacy is driving significant checkpoint inhibitor adoption, and analysts project the class alone could represent a \$29 billion global market by 2022. In short, checkpoint inhibitors are providing hope to metastatic patients who were previously considered to be on a path to palliative care.

Despite their performance, checkpoint therapies have a number of shortcomings. For instance, response rates are still only 20% to 30% on average (although there is significant variance in published response rates by tumor type), and response times can be prolonged, which can be an issue for patients with advanced metastatic disease.

Checkpoint inhibitors also come with significant side effects, especially when used in the combination regimens (e.g., with anti-CTLA-4) that are ubiquitous in the industry. (Also see "Combinations Continue To Drive Immuno-Oncology Deal-Making" - In Vivo, May 2017.) Checkpoint inhibitors' cost (at up to \$150,000 per year for monotherapy) represents a significant burden to health care systems and payers. Furthermore, prior exposure to checkpoint therapy may render patients ineligible for other immuno-oncology clinical trials.

Today, PD-1/PD-L1 immunohistochemistry tests (IHCs) are available as companion or complementary diagnostics for many approved indications, yet their predictive power is considered limited in many situations. As such, while these IHCs are broadly ordered by clinicians, decisions to use checkpoint therapy may often be based on a lack of therapeutic alternatives coupled with substantial patient demand. This is creating significant need for improved diagnostics to predict response to checkpoint therapy, monitor response and support sustained usage.

Complex Biology

Cancer pathways are already complex, and when you layer in the need to understand the host's immune system and potentially the microbiome, the complexity is greatly enhanced. Furthermore, given the inherent heterogeneity of both the tumor and immune cells, understanding the biology may be required down to the single-cell level.

Through a systematic review of clinical trials involving checkpoint inhibitor therapies, we identified over 1,200 ongoing or completed clinical trials involving checkpoint inhibitors going back to 2011. We then mapped the biomarker activity covered in those trial protocols to develop insights on pathways and biomarkers under exploration by biopharma and academic sponsors. Based on this mapping exercise, immunooncology biomarkers in solid tumors were broadly categorized into four areas:

1. Neo-antigen generation: Tumor cells generate neo-antigens as a result of genetic alterations; these neo-antigens are recognized by immune cells (via their T-cell receptor), resulting in immune-cell activation.

2. Immune activation: Recognition of tumor cell neo-antigens leads to immune-cell proliferation and pathway activation.

3. Immune evasion: Tumors have mechanisms to evade immune system attack, including evading recognition (cloaking), such as in the PD-1/PD-L1 checkpoint mechanism, and recruiting regulatory cells to suppress immune cell-killing ability.

4. Microbiome: There is also emerging evidence on the role of the microbiome in checkpoint therapy responsiveness, although this is in the very early stages.

Multiple pathways and biomarkers are being explored across categories. Neo-antigen generation trials are looking at TMB (tumor mutational burden), MSI (microsatellite instability) and DNA repair dysfunction. All three rely on the simple premise that



The industry needs to go through this current R&D wave to really understand which biomarker strategies work best by tumor type and indication.

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the more mutated or genetically unstable a tumor, the more foreign it will look to the immune system. TMB's role in checkpoint therapy responsiveness emerged quickly from collaborative tumor profiling efforts by academia and clinical laboratories such as **Foundation Medicine Inc.** and has since garnered significant investment and attention, most recently from the positive results of partner Bristol Myers Squibb's CheckMate 227 study. (Also see "Bristol's Opdivo/Yervoy Bid Will Show Whether Tumor Mutation Burden Is Ready For Prime Time" - Pink Sheet, February 5, 2018.)

Immune activation biomarkers are focused on measuring the presence of cancer-killing immune cells such as CD8+ T cells and the clonal expansion of those T cells, which suggests the immune system has recognized the cancer and is proliferating in preparation for tumor attack. Detection of killer immune cells has traditionally relied on immuno-phenotyping based largely on cell-surface markers, but recent activity also highlights interest in looking at HLA (human leukocyte antigen) genotypes and how they correlate with checkpoint therapy responsiveness. Expression of biomarkers correlated with immune pathway activation such as INF-gamma is also being explored as they are indicators of immune-mediated killing of cancer cells.

But cancers have cloaking mechanisms, and this drives the need to look at immune evasion biomarkers, which today focuses primarily on checkpoint activation. The current wave of cancer immunotherapies relies on the disruption of the PD-1/PD-L1 checkpoint interaction, and measuring PD-1 or PD-L1 expression is often correlated with inhibitor response. The industry is also beginning to look into the role of the microbiome in PD-1/PD-L1 inhibitor therapy responsiveness, as it may represent an important use case in microbiome-based diagnostics.

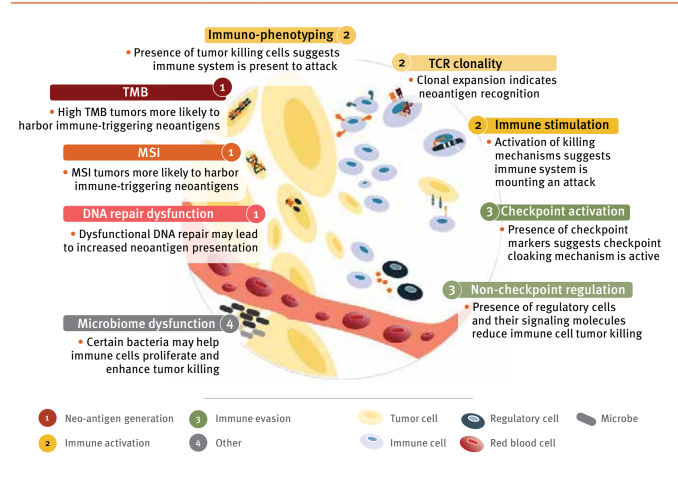
Clinical Trials Are Increasingly Assessing Multiple Biomarkers

The sheer growth in trial numbers is impressive, but there is another important trend to call out, which is the number of unique biological pathways/biomarkers being assessed per trial. (*See Exhibit 2.*)

Since 2015, there has been notable growth in trials measuring two or more

Exhibit 1

Overview Of Immuno-Oncology Pathways Under Exploration



pathways as part of the trial design, and this trend appears to be accelerating, with some trials looking at four or more pathways. It is important to note that while many trials may not specify a biomarker, many require tumor biospecimens as an enrollment criterion, suggesting that biomarkers may be explored and even submitted to regulators regardless of what is specified in the trial protocol. It is also important to mention that merely highlighting interest in exploring a particular pathway or biomarker as part of a clinical trial does not necessarily lock in a biomarker as part of the drug label.

Biomarkers Assessed Are Broadening Beyond PD-1

Checkpoint activation biomarkers (PD-1/ PD-L1) have seen the most activity and sustained growth, as they are directly related to the mechanism of action of checkpoint inhibitor therapies. However, exploration of other pathways and biomarkers is occurring in parallel. Trials looking at immune activation biomarkers, including immune stimulation, immunophenotyping (this includes tumor infiltrating lymphocyte [TIL] counting) and TCR clonality, have exploded since 2015 and are now being explored in 40% of biomarker-specified trials initiated in 2017.

Neo-antigen generation biomarkers have also grown significantly since 2015, with emphasis on MSI and TMB. MSI (for which Merck & Co. gained the industry's first biomarker-defined drug label) activity is the highest of the two, but it is also a more established biomarker, whereas TMB, which has gained prominence in the last year, is already being explored in registration trials by a number of biopharma players, including **Roche** and Bristol-Myers Squibb.

Since 2015, there has also been an uptick in assessing non-checkpoint regulatory biomarkers, such as IDO and FOXP3 (representing 8% of clinical trials initiated in 2017). And although still small today, there is a clear interest in assessing the microbiome's role in checkpoint therapy trials (<3% of trials initiated in 2017). (*See Exhibit 3.*)

Looking at biomarkers by development phase highlights that while checkpoint activation biomarkers are the most represented pathway in Phase III trials, most other pathways are being explored across development phases. (*See Exhibit 4.*)

Biomarkers Are Being Explored Across Tumor Types

Not surprisingly, overall trial activity is highest in NSCLC and melanoma, as these are indications where the first checkpoint therapy approvals were awarded. But there is also meaningful trial activity in other solid tumors such as breast, headand-neck, renal, colorectal (CRC), bladder and even liquid tumors. Interestingly, biomarkers are being explored across all tumor types, with CRC over-indexing relative to other tumors (due to the wellestablished link between MSI in CRC),

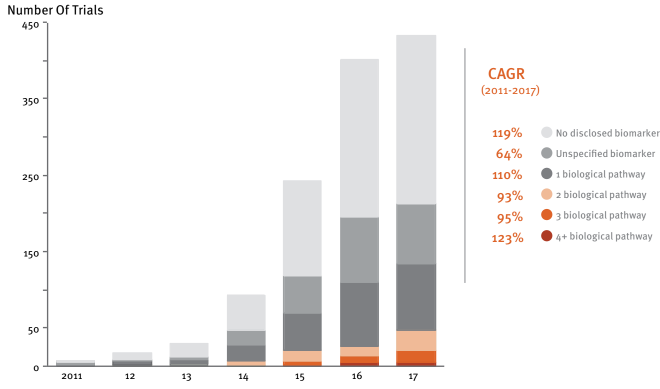
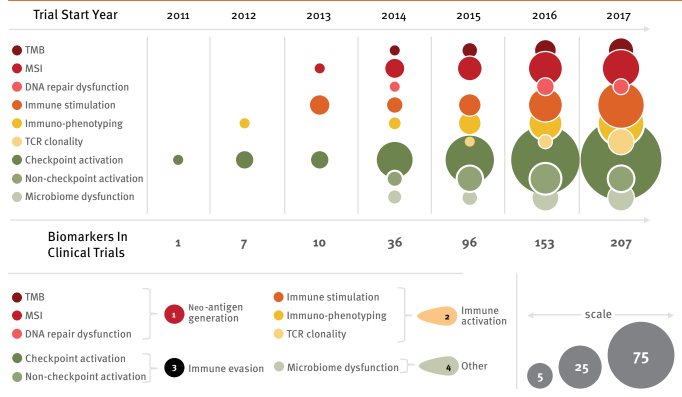


Exhibit 2 Number Of Biological Pathways Assessed In Checkpoint Inhibitor Clinical Trials

Note: 2, 3, and 4+ biological pathway CAGRs are from 2012-2017, and 2014-2017 respectively.

Exhibit 3

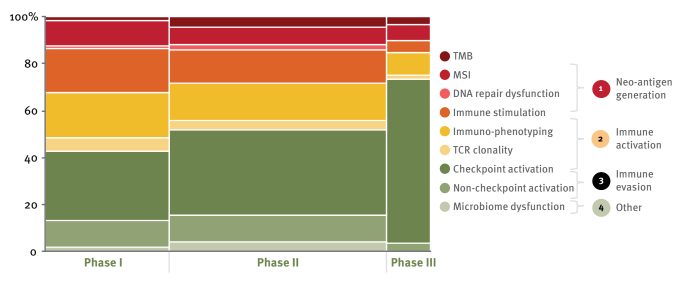
Biological Pathways Assessed In Checkpoint Inhibitor Clinical Trials By Trial Start Year



Notes: CAGRs may vary from start year of 2011-2014 but end Nov. 2017; does not include 13 retrospective studies without a known start date; includes double counting of trials if trial has more than 1 biological pathway interrogated.



Biological Pathways Assessed In Checkpoint Inhibitor Clinical Trials By Development Phase



Percent Of Biomarkers In Clinical Trials

Notes: Excludes 2 biomarker-specific studies, leveraging real-world evidence; includes double counting of trials if trial has more than 1 biological pathway interrogated.

whereas HCC (hepatocellular carcinoma), pancreatic and gastric cancers are underindexing on a relative basis.

Ultimately, with few exceptions, it appears that most tumors will benefit from biomarker analysis to predict checkpoint inhibitor responsiveness. That said, the industry needs to go through this current R&D wave to understand which biomarker strategies work best by tumor type and indication. (*See Exhibit 5.*)

A possible future outcome may be that many pathways and biomarkers will need to be assessed regardless of tumor type, to tailor IO therapy to patients by looking at the underlying biology of the tumor, the host immune system and even the microbiome.

Multiparameter Diagnostic Modalities Will Be Critical

Given the number and types of biomarkers explored (DNA, RNA, protein), multiple diagnostic modalities are being employed, including next-generation sequencing (NGS), quantitative polymerase chain reaction (qPCR), **NanoString Technologies Inc.**'s *nCounter*, IHC and flow cytometry. NGS (covering both DNA and RNA sequencing) appears to be poised to address the majority of IO biology and pathways, and its increasing prominence in tumor profiling in metastatic disease could position the technology as a front-runner in IO diagnostics. However, high-parameter flow cytometry and IHC are also expected to be important IO diagnostic tools in the long run given their ability to detect expressed proteins at the single-cell level. (*See Exhibit 6.*)

Another notable point in the IO diagnostics space is the emergence of multi-parameter RNA expression and single-cell NGS. Multi-parameter expression analysis has the potential to include multiple areas of IO biology and pathways, essentially covering both immune activation and evasion. And because both tumors and the immune system are marked by their cellular heterogeneity (due to the genetic instability of tumors and the adaptability of immune cells to new threats), the need for single-cell NGS is also gaining prominence in research.

Although the cost to assess multiple biomarkers to predict checkpoint responsiveness may represent a step-change from the single biomarker companion diagnostics associated with most targeted therapies, the clinical and economic need associated with tailoring the use of checkpoint inhibitors will likely support this added cost.

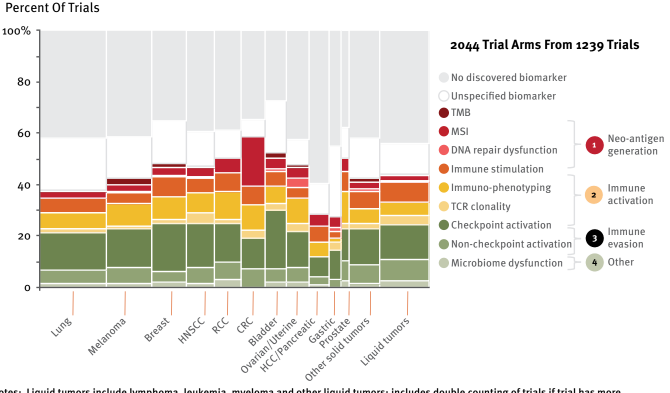
It is important to note that many diagnostic enablers, including platform companies such as **Illumina Inc.**, **Thermo Fisher Scientific Inc.**, **HTG Molecular Di**- agnostics Inc. and NanoString, as well as clinical laboratories such as Foundation Medicine, OmniSeq LLC and Caris Life Sciences, are developing immuno-oncology diagnostic solutions that span multiple biological pathways and are actively partnering with pharma. (See Exhibit 7 online.) OmniSeq, a clinical laboratory associated with Roswell Park Cancer Institute, now offers Immune Report Card, which looks at pathways across neo-antigen generation, immune activation and immune evasion. A recent commercial partnership between OmniSeq and Laboratory Corp. of America Holdings has the potential to significantly expand access to OmniSeq's Immune Report Card by leveraging Lab-Corp's extensive channel.

NGS And Big Data Are Accelerating The Biomarker Innovation Cycle

Against this backdrop, there is an acceleration in the biomarker innovation cycle driven by the comprehensive genomic profiling of tumors using NGS. LEK Consulting estimates that 10% to 20% of metastatic patients receive NGS profiling in the US today, and that number could increase two to three times in the next three to five years (which represents hundreds of thousands of patient cases annually).

Many of the NGS providers are building large data sets incorporating genomics,





Notes: Liquid tumors include lymphoma, leukemia, myeloma and other liquid tumors; includes double counting of trials if trial has more than 1 biological pathway interrogated.

clinical and outcomes data which is enabling identification and retrospective validation of new biomarker associations directly from real-world clinical cases. This activity has the potential to validate new biomarker associations backed by a significant number of patient cases (often dwarfing what is feasible with traditional clinical trials). Supported by this strength of evidence, it is expected that guidelines and clinical adoption of these biomarkers will follow quickly. Conceptually, this represents the application of big data in precision genomic medicine.

However, as highlighted earlier, this is no longer purely conceptual. The rapid rise of TMB represents a paradigm shift. TMB isn't even a definable genomic biomarker in the traditional sense. TMB is affiliated not with a specific gene or pathway, but rather with an observation that tumors with a relatively high frequency of mutations (high TMB) tend to respond better to checkpoint inhibitors (thought to be due to increased neo-antigen presentation). Interestingly, TMB also represents a new situation where diagnostics companies have forged the path and biopharma has followed suit. This stands in significant contrast to other targeted therapies where biomarkers and their associated complementary/companion diagnostics were largely validated by biopharma-sponsored pivotal trials.

TMB is also creating a virtuous discovery cycle that may only continue churning out new and more nuanced biomarker associations. NGS is required to measure TMB, and as TMB becomes more routinely adopted it is an important driver to continued NGS adoption. Furthermore, NGS panel sizes continue to grow, and digitization of health care data by players such as **Flatiron Health Inc.** and **COTA Inc.** is enabling the creation of massive minable data sets.

The Road Ahead

There is a long road ahead for the IO diagnostics space to mature. As discussed earlier, the science and supporting evidence need to be developed, and there are many directions in which they could go. Will it end with a reductionist biomarker strategy reliant on a few highly predictive biomarkers, or will it end with a more comprehensive biomarker strategy that looks at many pathways in concert? Certainly, the biology of IO therapies, which relies on the interaction between a dynamic tumor and the immune system suggests the more comprehensive strategy may prevail in the long run, but it will take time for that to materialize.

Separately, the industry needs to drive standardization, not only in pathways and biomarkers assessed, but also in defining standards around interpretation, including defining thresholds for what constitutes a biomarker-high or biomarker-positive result. Standardization and concordance across sample types assessed will also need to be worked out, including understanding cell heterogeneity (for both tumor and immune cells) and between tissue-based samples and discriminating between those derived from biofluids including CTCs, peripheral immune cells, cell-free components, exosomes and so forth.

The industry will also ultimately want to consolidate testing into a standard set and will not support a different test for every therapeutic option under consideration. This means there will need to be cooperation and collaboration across the industry,

Exhibit 6 Overview Of Biomarkers And Modalities In IO Diagnostics

Biological pathway		System	Analyte		Example biomarkers	Key diagnostic modalities	
1. Neoantigen generation	тмв	Tumor cell	Somatic mutation density	DNA	тмв	NGS	
	MSI		Short tandem repeats		Mismatch-repair status	NGS qPCR	
	DNA repair dysfunction		Mutations in DNA repair gene		HRRm, POLE GA	NGS qPCR	
2. Immune activation	Immune stimulation	Immune cell	Expression of immune cell activation markers	RNA protein	IFNg, TNFa, CRP, cytokines	NGS qPCR nCounter	IHC Flow
	lmmuno- phenotyping				CD8, CD4, CD3, CD25, HLA	NGS qPCR nCounter	IHC Flow
	TCR clonality		Clonal population analysis of variable region	DNA	TCR regions	NGS	
3. Immune evasion	Checkpoint activation	Tumor and Immune cell	Expression of checkpoint immune evasion markers	RNA protein	<u>TC:</u> PD-L1, PD-L2, TAA, Ki67 <u>IC:</u> CTLA-4, PD-1, LAG-3	NGS qPCR nCounter	IHC Flow
	Non- checkpoint regulation				TC: IDO IC: FOXP3	NGS qPCR nCounter	IHC Flow
4. Other	Microbiome dysfunction	GI/stool	Microbiome composition and genetic diversity	rRNA	165 rRNA	NGS	

SOURCE FOR ALL EXHIBITS: LEK Consulting

and looking at other diagnostics markets, a reasonable assumption suggests three to five major competitors will emerge as leaders.

Access to novel diagnostic approaches will also need to be addressed. Today, many of the emerging IO diagnostic tests require specialized instrumentation and operators, may take many weeks to process and are not well-reimbursed. Clearly, as the industry scales, this will need to change and fit better into therapeutic decision-making windows and health-care economics.

Biopharmaceutical companies will need to adapt in this new fast-paced biomarker environment. Trying to stay abreast of the biomarker landscape in a sequential fashion with each new biomarker approval may put companies into endless catch-up mode.

An alternative strategy could be to proactively assess multiple pathways and biomarkers at the outset, leveraging basket trials with fast conditional approvals to match therapeutic options to each patient's unique (and ever changing) biomarkers.

Organizationally, biopharmaceutical

companies will need to change across many functions. Deeper embedding of biomarker/diagnostics groups into development and commercial functions will be required. Business development activity with diagnostics partners will need to focus on broader collaborations with a focused set of diagnostic companies, with the ultimate goal of creating industry-standard diagnostic solutions.

Market access and pricing functions will also need to consider how evolutionary biomarker strategies impact pricing and reimbursement of both the IO diagnostics and therapeutics.

Regulators will do well to enable such forward-thinking approaches. Just like the immune system, the industry needs to adapt to enable IO companion therapies and diagnostics to reach their full potential. IV005261

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READ MORE ONLINE

Who are the major players in immuno -oncology diagnostics? See our online exhibit of the most active solutions developers and partnering prospects.

Follow this link to our list of the leading companies in IO diagnostics.

http://bit.ly/2EHLnPg

If you have questions about the article or would like more information, please contact Alex Vadas at a.vadas@lek.com



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