BIOMARKERS AND DRUG DEVELOPMENT: REGULATORY PERSPECTIVE

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• Views expressed in this presentation are those of the speaker and do not necessarily represent an official FDA position.

• I do not have any financial disclosures regarding pharmaceutical drug products.
OVERVIEW

• Communicating from a common understanding: BEST resource
• Biomarker Integration into Drug Development
• Biomarker Development and Qualification
• Biomarker-related resources
• Summary
**Definition**: a defined characteristic that is measured as an 1) indicator of normal or pathogenic biological processes or 2) response to an intervention.

Broadly defined, with multiple biomarker types including molecular, histologic, radiographic, and physiologic. (i.e., serum protein, change in tumor size by imaging study, algorithm for QT determination on ECG)

Characteristic is not a *clinical* assessment of how a patient feels, functions, or survives (contrasted with Clinical Outcome Assessments or COAs)

Although a biomarker may be used by clinical or basic science research communities, regulatory acceptance focuses on a drug development context that is supported by data for that context. Considerations include:

- Reproducibility of data (e.g., high rate of discordant conclusions RE biomarkers in the published literature)
- Adequacy of the analytic device to assess biomarker’s reliability
- Feasibility of the biomarker should a drug be approved (e.g., will the analytic be widely available and capable of integration into clinical practice paradigms)
BEST: BIOMARKERS, ENDPOINTS, AND OTHER TOOLS RESOURCE

- A glossary of terminology and uses of biomarkers and endpoints in basic biomedical research, medical product development, and clinical care
- Created by the NIH-FDA Biomarker Working Group
Biomarker Classes from a Drug Perspective

- **Susceptibility/Risk**: Indicates potential for developing disease or medical condition in an individual who does not currently have clinically apparent disease or the medical condition.

- **Diagnostic**: Detects or confirms the presence of a disease or condition of interest or to identify individuals with a subset of the disease.

- **Monitoring**: Assesses status, through serial measurement, of a disease or medical condition including degree or extent of disease.

- **Prognostic**: Identifies likelihood of a clinical event, disease recurrence or progression, in patients who have the disease or medical condition of interest in the absence of a therapeutic intervention.

- **Predictive**: Identifies patients who are more likely to experience a favorable or unfavorable effect from a specific treatment.

- **Pharmacodynamic/Response**: Indicates that a biological response has occurred in a patient who has received a therapeutic intervention. May become clinical trial endpoints and for a very small subset, surrogate endpoints.

- **Safety**: Indicates the likelihood, presence, or extent of toxicity to a therapeutic intervention when measured before or after that intervention.
“Fit for Purpose”: BEST Biomarker Classes in Perspective -- Match Biomarker to a Drug Development Goal and Data-supported Relationship

- **“Normal” Physiology**
  - Susceptibility/Risk
  - **Pathologic Changes**
    - Descriptive
      - Time progression
      - Key factors / events
  - **Altered Physiology**
    - Descriptive
      - Threshold of concern
  - **Clinical Disease**
    - Diagnostic
    - Monitoring
    - Prognostic
- **Improved Clinical Benefit**
  - Surrogate Endpoint
  - **Non-Progression Or Reversal**
    - Response
  - **Change in Physiology**
    - Pharmacodynamic
    - Predictive
    - Safety
  - **Therapeutic Intervention**
Each of these elements share importance to drug approval. Since any element can lead to failure, important to optimize as appropriate and feasible.
CONSIDERATIONS FOR BIOMARKER UTILITY

Context of Use (COU): 1) BEST biomarker category and 2) how the biomarker impacts the clinical trial or drug development program

What question is the biomarker intended to address. Examples include:

- Inclusion/exclusion criteria for prognostic or predictive enrichment?
- Alter treatment allocation based on biomarker status?
- Result in cessation of a patient’s participation in a clinical trial because of safety concern?
- Result in adaptation of the clinical trial design?
- Establish proof of concept for patient population of interest?
- Support clinical dose selection for first in human or Phase 3 studies?
- Evaluate treatment response (e.g. pharmacodynamic effect)?
- Support regulatory acceptability of a surrogate endpoint for accelerated or traditional approval?

“Total Kidney Volume, measured at baseline, is a prognostic enrichment biomarker to select patients with ADPKD at high risk for a progressive decline in renal function (defined as a confirmed 30% decline in the patient’s estimated glomerular filtration rate (eGFR)) for inclusion in interventional clinical trials. This biomarker may be used in combination with the patient’s age and baseline eGFR as an enrichment factor in these trials.”

Note: These pathways do not exist in isolation and many times parallel efforts are underway within or between pathways. All share common core concepts, are data-driven, and involve regulatory assessment and outcomes based on the available data.
SCIENTIFIC COMMUNITY CONSENSUS APPROACH FOR BIOMARKER DEVELOPMENT

Strengths
- Extensive knowledge base of exploratory biomarker data in published literature
- Opportunity for broad and multiple community inputs
- Public access and cost-sharing approach (e.g., NIH and other grant funded research)

Limitations
- Published data may not be not reproducible
- Protracted time for consensus building
- Variability of study designs, populations, and analytics
- Applicability to regulatory paradigms
DRUG APPROVAL (IND/NDA/BLA)
APPROACH FOR BIOMARKER DEVELOPMENT

**Strengths**
- Generally, biomarker use has a well-defined purpose
- Data (clinical trial information) available to the biomarker developer
- Opportunities to bring in outside experts
- Company retains marketing advantage

**Limitations**
- Biomarker may not be generalizable
- Limited opportunities for additional data sources
- Company responsible for development costs
- Limited opportunities for engagement with outside stakeholder groups
- Biomarker information in drug labels and reviews are available only upon drug approval

Drug Approval Process

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BIOMARKER QUALIFICATION APPROACH FOR BIOMARKER DEVELOPMENT

**Strengths**
- Context of use clearly established
- Opportunity to pool resources, share costs and bring outside experts
- Leverage outside stakeholder groups
- Outcome is a public guidance with supporting reviews

**Limitations**
- If part of a group effort, stakeholders may have differing goals, level of commitment, and engagement
- Data (clinical trial information) may not be readily available
- Data sharing and aggregation may be challenging

SOME ENABLERS FOR BIOMARKER DEVELOPMENT

• Data standards (e.g., CDISC efforts)
• Data quality
• Data reproducibility
• Data sharing
• Assay/imaging pre-analytic standardization
• Assay/imaging protocols/SOPs
• Evaluating impact on clinical trial elements (e.g., choice of cut-point on number of patients screened vs enrolled)
CONCEPTUAL FRAMEWORK FOR BIOMARKER DEVELOPMENT FOR REGULATORY ACCEPTANCE

In Drug Development
- Need Statement
  - Class of Biomarker.
  - What is the question the biomarker is addressing.
- COU
  - Evaluate Compared to Status Quo
- Benefit
  - Improved sensitivity
  - Improved selectivity
  - Mechanistic context
- Risk
  - Consequence of false positive
  - Consequence of false negative
- To Patient
  - Informs Required Stringency of EC

Evidentiary Criteria
- Characterization of Relationship Between the Biomarker and Clinical Outcome
- Biological Rationale for Use of Biomarker (if Known)
- Type of Data and Study Design (i.e. Prospective, Retrospective, etc.)
- Independent Data Sets for Qualification
- Comparison to current standard
- Assay performance
- Statistical Methods to Use
The Specific Context of Use for a Biomarker Drives the Extent of Evidence Needed for Qualification

**Analytical Validation**
- Reference Ranges/Decision Points
- Pre-Analytical and Assay Performance Characteristics
- Analytical Rigor/Reproducibility
- Sample Handling/Stability

**Clinical Validation**
- Study Design Acceptability
- Clinical Meaningfulness/Decision Points
- Benefit/Risk Assessment

**ANALYTICAL ASSAY AND CLINICAL VALIDATION CONSIDERATIONS IN BIOMARKER QUALIFICATION**
21st Century Cures legislation: Section 507 Qualification of Drug Development Tools

- 21st Century Cures and PDUFA VI increasingly places FDA as an active participant in drug development, broadening our traditional regulatory role
- Formalizes a three-step submission process
  - Letter of Intent
  - **Qualification Plan**
  - Full Qualification Package
- A transparent process – so all stakeholders are aware of tools in development, stage, and FDA determinations/recommendations
- Requires setting and implementing “reasonable timeframes” for submission review/decision
TRANSPARENCY PROVISIONS

Under 21CC, DDT qualification becomes a transparent public process:

- All interested parties know what tools are in development, stage of development, and FDA determinations including rationale
- Information about the submission and FDA’s determination including recommendations will be posted on DDT website
- For legacy projects, we plan to post only new information after transition (e.g., we will not make public information prior to legislation enactment or to agreement to transition to 507)
- De facto Letter of Support (LOS) as part of DDT engagement
Acceptance decision for each submission (LOI, QP, FQP) based upon scientific merit:

- Does the proposal address an impactful drug development need?
- Is there enough information to suggest a likelihood of success?
- What is the feasibility of the proposed analytical biomarker measurement approach?

Prioritization of review of submissions based upon:

- “the severity, rarity, or prevalence of the disease or condition targeted by the drug development tool and the availability or lack of alternative treatments for such disease or condition; and
- the identification by the Secretary or by biomedical research consortia and other expert stakeholders, of such drug development tool and its proposed context of use as a public health priority” (italics added)
CONTENT FOCUS FOR SUBMISSION TYPES

• LOI Submission: Feasibility assessment of proposal will include information to support that measurement of the novel DDT is, in fact, possible.

• QP Submission: Project development plan from concept to information to be developed/provided to support the DDT’s COU. For biomarkers, to determine clinical utility and clinical validation, important to know that the analytical validation has been completed and information submitted to QP.

• FQP Submission: Review of data to support the clinical validation of the DDT for the COU
COMPOSITE BIOMARKERS CONSIDERATIONS

Terminology: Composite, panel, multi-modal, score, etc.

- As a start, list out the individual biomarker components
- The “final” list can evolve but plan accordingly

Strategies:

- Start with a single COU, explore biomarkers/measurement methodologies individually and then build into composite once value has been demonstrated
- Alternatively, again with a single COU, start with composite of most promising candidates and then refine

Measurement Scenarios:

- Each member of the composite is measure with a separate platform (each independently validated). The readouts are then “manually” transformed into a composite
  - Single composite readout/interpretation = cut points (ex. Mild, Moderate, Severe)
  - Group assessment. Still have cut points for “biomarker positivity” but interpretation simplified (ex., if any 2/7 positive, then composite is “positive”)
- The individual biomarkers of the composite are measured by a single device
  - Does the device readout individual values (ex. Chem7) or combine (ex. Score)?
  - If score, then the algorithm which generates the score needs to be provided
THREE-TIERED INTERNAL REVIEW

• DDT Program Assessment and Recommendations
  • Work with requestor to clarify DDT, COU, and project proposal
  • Provide tool-specific recommendations based on past and ongoing projects

• Discipline-specific SME Assessment and Recommendations
  • Includes OND division management participation
  • Evaluate based on regulatory precedent, current disease-specific challenges, and level of impact on drug development programs

• CDER DDT Committee Assessment, Recommendations, and Decision
  • Opportunity for broad senior CDER input early and throughout in the process
  • Work towards greater consistency across therapeutic areas and divisions
BIOMARKER QUALIFICATION PROGRAM (BQP) UPDATES

• Legacy projects fully transitioned to 507 process
• Coordination with CBER for inter-Center consistency
• Continued collaboration with EMA for alignment on process and approach
• LOI, QP, and FQP submission content outlines now developed and posted
• In planning
  • New IT platform (submission portal, document knowledge management, business tools)
  • Guidances for DDT qualification process, BQ evidentiary framework, surrogate endpoints, and BQ analytical validation
• Public posting of information for transparency
TABLE OF SURROGATE ENDPOINTS

21st Century Cures Act, Subtitle B—Advancing New Drug Therapies

SEC. 507. QUALIFICATION OF DRUG DEVELOPMENT TOOLS.

“Transparency

“(E) A comprehensive list of—

“(ii) all surrogate endpoints which were the basis of approval or licensure (as applicable) of a drug or biological product (including in accordance with section 506(c)) under section 505 of this Act or section 351 of the Public Health Service Act.”

- 101 adult and 56 pediatric disease/patient population/surrogate endpoint combinations
- 12 surrogate endpoints that may be appropriate for use in drug approval even though no successful drug program as of yet
- More disease/therapeutic areas use surrogates than commonly discussed
- Will be updated every 6 months
IND TYPE C MEETING FOR NOVEL SURROGATE ENDPOINTS

- PDUFA VI Commitment
- Meeting package due at time of request that includes preliminary human data indicating drug has an impact on the SE at a dose that is “generally tolerable”

Package content examples include:
- Rationale for use of surrogate endpoint (SE)
- Relationship of SE with casual pathway(s)
- Threshold for change required to demonstrate clinical relevance
- Consistency of SE response
- Reliability of quantifying changes in clinical outcome before and after tx
- Predictive value of therapeutic-induced changes in SE
- Off-target effects of therapy
- Reliability of measurement tool to detect SE
THANK YOU FOR YOUR ATTENTION

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