Issues in Clinical Trial Design for Companion Diagnostic Devices

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Presentation Outline

- Overview of Regulation of In Vitro Diagnostic Devices (IVDs)
- Scope of Data to Support Approval/Clearance of an IVD Device
- Companion Diagnostics Co-development: Benefits and Challenges
- Useful Tools
Definition of an In Vitro Diagnostic Device

“Reagents, instruments, and systems intended for use in the diagnosis of disease or other conditions, including a determination of the state of health, in order to cure, mitigate, treat, or prevent disease or its sequelae. … for use in the collection, preparation, and examination of specimens from the human body.”

[21 CFR 809.3]
Sources of IVD Devices

Who is the innovator of the IVD?

IVD Manufacturer
- FDA Cleared/Approved Kits
- RUOs and IUOs
- IVD MIA

Clinical Laboratory
- Contract Manufacturing
- ASRs
- Non-ASR LDTs
Regulatory Authority

- Federal Food, Drug and Cosmetic Act
  - Established regulatory controls for Medical Devices (May 28, 1976)

- Code of Federal Regulations, Title 21, Part 800
Human Subject Regulations

Definition of a “Human Subject”

– Human who participates in research either as a recipient of the test article or as a control. A healthy human or a patient

– Subject is an individual on whom or on whose specimen an investigational device is used
FDA Human Subject Protection Regulations

- **21 CFR Part 50**: Informed consent and limited emergency exceptions

- **21 CFR Part 56**: IRB review

- **21 CFR 812**: Disqualification of an Investigator (812.119)

  - Apply to all FDA clinical investigations
Guidance for Sponsors, IRBs, Clinical Investigators and FDA Staff

“Guidance on Informed Consent for In Vitro Diagnostic Device Studies Using Leftover Human Specimens that are Not Individually Identifiable”
(4/25/2006)

http://www.fda.gov/cdrh/oivd/guidance/1588.html
CDRH Total Product Life Cycle Regulatory Approach

Postmarket

Design

Analytical Evaluation

Quality Systems 21CFR §820

Clinical Evaluation

Postmarket Surveillance

Premarket

Compliance

Approval/Clearance

http://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfTPLC/tplc.cfm
Pre-Market Risk Based Regulation

Class I and II

Knowledge Mitigates Risk

Class I - Low likelihood of harm
Register & list
General Controls. 510(k)

Class II - Moderate likelihood of harm
or risk can be mitigated
Special Controls. 510(k)

Class III - High or unknown
likelihood of harm
Significant Risk
Pre-Market Approval
Basis of Pre-Market Device Review: Safety and Effectiveness

• Safety
  – Are there reasonable assurances, based on valid scientific evidence that probable benefits to health from use of the device outweigh any probable risks? [860.7(d)(1)]

• Effectiveness
  – Is there reasonable assurance based on valid scientific evidence that the use of the device in the target population will provide clinically significant results? [860.7(e)(1)]
Intended Use of the IVD

“Intended Use”-driving force of the scientific review

- Understanding:
  Integration of disease(s)/condition(s).
  Integration of patient clinical management and public health (surveillance)

  - **Who** will be tested, where and when: outpatients, inpatients, pediatrics, adults, acutely ill, etc.

  - **What** are the appropriate specimens: timing, handling

  - **How** test result(s) may be used: patient management
Scientific Review: 
Device Performance

- **Analytical** Performance Characteristics
  Reliability and accuracy of analyte measurements

- **Clinical** Performance Characteristics
  Clinical sensitivity and specificity
  Positive and negative predictive values

- **Labeling**
  Intended use, device design, directions for use, warnings/limitations, result interpretation, performance
Demonstrating Evidence for Safety: Analytical Studies

• Likelihood of false positives?
  – Cross-reactivity and other interferences
  – Carryover and contamination

• Likelihood of false negatives?
  – Limits of detection
  – Matrix effects
  – Interference
Demonstrating Evidence for Effectiveness: Clinical Studies

- Well-controlled clinical evaluations:
  - Clinical plan and protocol
  - Defined objective(s) and methods

- A test device with standardized design and performance

- Other evidence: case histories, literature, reproducibility etc. where appropriate
FDA Guidance Relevant to In Vitro Diagnostic Device Clinical Trials

- “In Vitro Diagnostic (IVD) Device Studies-Frequently Asked Questions”
  Published 10/25/07
Companion Diagnostic Policy Emerges..

- Healthcare practitioners now rely on information from companion diagnostic devices to help make critical treatment decisions.

- Companion diagnostics
  - Provide benefit in optimizing treatment
  - Bring risk if test result is incorrect
What is a Companion Diagnostic?

- An *IVD companion diagnostic device* is an in vitro diagnostic device that provides information that is essential for the safe and effective use of a corresponding therapeutic product.

  - *If the safe and effective use of the therapeutic product requires a particular test result, that test is a companion diagnostic*
Draft Guidance Published

- “Draft Guidance for Industry and FDA Staff; In Vitro Companion Diagnostic Devices” published July 14, 2011
  - Docket FDA-2011-D-0215
  - Comment period extended to October 12, 2011
  - Final guidance still pending publication
Regulatory Concept: Premarket Review

- Premarket review and clearance or approval of the companion diagnostic will typically be required prior to, or contemporaneously, with approval of the therapeutic product
  - Assurance that the diagnostic has been appropriately validated for its intended use
  - Risk-based regulation still applies
Regulatory Concept: Co-approval

- Companion Dx and therapeutic product depend on each other
- Co-approval required*
- Failure/lack of test approval = no therapeutic product approval

* Exceptions will exist
What Types of Tests Could Be Companion Dx?

- Identify patients likely to respond or not respond to a particular medical product

- Predictive test that can select either a population that is likely to respond, or a population that is not likely to respond

  - Generally validated in Phase 3 trials, although could be a post-approval addition, e.g., KRAS
What Types of Tests Could Be Companion Dx? (2)

- Identify subgroups of the larger population with poor prognosis who are likely to benefit from a particular therapeutic product.

- Test selects those who might benefit from treatment with a therapeutic product due simply to otherwise poor prognosis.
What Types of Tests Could Be Companion Dx? (3)

• Identify patients likely to be at increased risk for serious adverse reactions as a result of treatment with a particular therapeutic product (predictive)

• Could be a test to show patient likely to, or unlikely to suffer SAR
  • Validation design can be tricky
  • Consideration of whether other therapies are available, seriousness of disease to be treated
What Types of Tests Could Be Companion Dx? (4)

• Monitor response to treatment for the purpose of adjusting treatment (schedule, dose, etc) to achieve improved safety or efficacy
  – Specific test performance needed, specific test values critical
  – Not generally accepted biomarkers for status
What Types of Tests Could Be Companion Dx? (5)

- *Individualize the dose of particular therapeutic product*

- Predicts safe/effective dose based on specific test result
What Types of Tests Could Be Companion Dx? (6)

- Use as integral part of therapeutic clinical trials conducted to support market approval of a therapeutic product e.g. selection of trial participants

- Primary trial analysis performed using diagnostic device data to demonstrate therapeutic performance
  - If test was used in any way to define trial success, it will need to be available to select the same population when therapeutic is approved
### Examples

FDA cleared / approved device in reference to the drug
FDA drug package insert sites the relevant test

<table>
<thead>
<tr>
<th>Efficacy Test</th>
<th>Safety Test</th>
<th>Drug</th>
</tr>
</thead>
<tbody>
<tr>
<td>HER-2/neu IHC, FISH &amp; CISH</td>
<td></td>
<td>Trastuzumab (Herceptin)</td>
</tr>
<tr>
<td>c-kit IHC</td>
<td></td>
<td>Imatinib mesylate (Gleevec)</td>
</tr>
<tr>
<td>EGFR IHC</td>
<td></td>
<td>Cetuximab, Panitumumab</td>
</tr>
<tr>
<td>BCR-ABL or t(9:22)</td>
<td></td>
<td>Imatinib mesylate (Gleevec)</td>
</tr>
<tr>
<td>EGFR mutations</td>
<td></td>
<td>Gefitinib, Erlotinib</td>
</tr>
<tr>
<td>UGT1A1 mutations</td>
<td></td>
<td>Irinotecan</td>
</tr>
<tr>
<td>TPMT</td>
<td></td>
<td>6MP &amp; Azathioprine</td>
</tr>
<tr>
<td>CYP450 (2D6) mutations</td>
<td></td>
<td>5-HT3R antagonist, codeine derivative</td>
</tr>
<tr>
<td>CYP2C9 &amp; VKORC1</td>
<td></td>
<td>Warfarin</td>
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Co-development Questions Pertaining to both Diagnostic and Drug

• Diagnostic Use
  – Does the test make a difference?
  – What if the test parameters (or the test itself) changes during or after development?

• Drug Development Program
  – Acceptable sequential or single adaptive trial designs?
  – Evidentiary standards
  – Generalizability, multiplicity, and other problems
Co-development Benefits

Evaluate drug and device in one trial

- For pharmaceutical companies
  - potential for optimum patient population and smaller future trials
  - improved drug effect if marker effective

- For diagnostic companies:
  - new type of diagnostic claim
  - well characterized subjects
  - extensive follow-up
Drug and Device Pathways from Pre-Clinical to Launch: Ideal Time Lines

**Drug Developmental Pathway**

- Basic Research
- Prototype Design or Discovery
- Preclinical Development
- Clinical Development
  - Phase 1
  - Phase 2
  - Phase 3
- FDA Filing Approval Launch

**Device Developmental Pathway**

- Basic Research
- Analytical Validation
- Feasibility Analysis
- Clinical Validation
- Qualification of Biomarker
- FDA Filing Approval Launch
Challenges of Co-development (1)

- May not be adequate data early on to determine the best biomarker to measure; whether test needed
- Appropriate statistics (may allow for e.g., adaptive trial design if drug not effective in the general population)
- Test used in drug trials not the marketed version (platform change)
- Appropriate storage of clinical trial samples; IRB, IC
Challenges of Co-development (2)

- Test analytical performance not validated prior to use in clinical trial

- Multiple tests (with different performance?) used to enroll subjects in clinical trial

- Bridging studies from CTA to companion Dx need high sample ascertainment (> 90% recommended)

- No way to determine test performance if only marker + samples available
  - Results in selection, but not predictive, claim
Challenges of Co-development (3)

Enrolment Issues:

- If test is part of inclusion/exclusion criteria for therapeutic trial
  - Use a central site to do ALL testing (not confirmation of local test)
  or:
  - Use SAME test (same demonstrated performance characteristics) at ALL sites
and:
  - Assure any test used has adequately validated measurement characteristics (analytical validation)
Analytical Validation

- Use a test for patient selection/stratification that has been as completely analytically validated as possible, with acceptable performance characteristics.

- For tests that detect multiple possible genetic changes, e.g., multiple mutations within a gene, should be analytically validated for each change to be detected.

- When performing analytical validation, consideration should be made of the ultimate specimen source to be used once drug is on the market, i.e., FFPE tissue, blood, CSF.
Testing Protocol/Samples

- Neither the test nor the testing protocol should be altered once the pivotal trial is begun.

- Sponsors should save samples from all patients enrolled in the trial. When only test-positive patients are enrolled:
  - do not prescreen for eligibility with a different test
  - save samples from patients who are test-negative and are thus not enrolled in the trial

- If a change in specimen type is anticipated for future testing, obtain paired samples on initial testing.

- For any samples that have been stored prior to testing, perform validations to assure that analyte is stable under storage conditions.
Policy and Guidance

• Schedule pre-Sub. meetings with CDRH as soon as test identified to discuss design of clinical studies etc. Helpful if both diagnostic and pharmaceutical reps. present

• Alert lead FDA center that therapy application includes a diagnostic device

• Companion diagnostics require compliance with FDA medical device regulations, regardless of test manufacturer

• A diagnostic test approval/clearance needed prior to drug approval in most cases

• Final intended use of test often depends on outcome of therapeutic clinical trial
Clinical Validation of Companion Dx

- Companion Dx (or its prototype or CTA) used within pivotal therapeutic trial
  - No systematic difference whether normal or accelerated approval of therapeutic

- Analysis of diagnostic test device performance related to therapeutic trial outcome
  - Therapeutic trial fails = diagnostic test not informative
  - Therapeutic trial succeeds = diagnostic (generally) informative
    - Diagnostic test label will reflect use in pivotal trial
  - Generally, no additional clinical validation of diagnostic needed
    - For predictive claims, therapeutic trial should be powered to detect differences in response by diagnostically defined strata.
Relabeling of a Therapeutic

- When new evidence becomes available to address a serious safety issue for an approved therapy
  - Relabeling of the therapy to include a test may be appropriate
    - **Required** test: test must usually be approved and available at time of relabeling
    - **Recommended** test: test ideally to be approved, but label change will not be delayed for approval
Conclusions: Co-development

- Drug-device trial allows for evaluation of broader range of claims

- Many complex issues in designing studies

- Use of unapproved /cleared IVDs or off label use of unapproved/cleared tests is an issue

- Use CDRH pre-Sub. process and CDER pre-IND process when planning study design and execution

- More research and collaboration between Diagnostic and Drug companies is vital
Tools: Standards for Evaluating Diagnostics

• Medical Devices Standards Database

• Clinical Laboratory Standards Institute (CLSI)
  – Develop global consensus standards and guidelines for healthcare testing (industry, government, professional)
  – Evaluation Protocols (EP) for study design/analysis
  – www.clsi.org

• ISO (International Standards Organization)
  – Standards for estimating bias and imprecision of test methods
Selection of CLSI Documents

- C28-A3. Reference Intervals
- EP5-A2. Evaluation of Precision Performance of Quantitative Measurement Methods
- EP17-A. Limits of Detection and Limits of Quantitation
- EP21-A. Total Analytical Error
- GP10-A. Assessment of the Clinical Accuracy of Laboratory Tests Using Receiver Operating Characteristic (ROC) Plots;
- MM17-A. Multiplex Nucleic Acid Assays
Information: CDRH
Homepage - Diagnostics

www.fda.gov/cdrh

- Device Classification Database
- Device Advice
  - http://www.fda.gov/cdrh/devadvice
- Register for “What’s New”
- Guidance Documents
- IDE Information
- Much more...
Summary of Presentation

• Covered Overview of Regulation of In Vitro Diagnostic Devices (IVD)

• Covered Scope of Data to Support Approval/Clearance of an IVD Device

• Discussed the Benefits and Challenges of Co-development for Companion Diagnostics

• Useful Tools
Questions?

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