



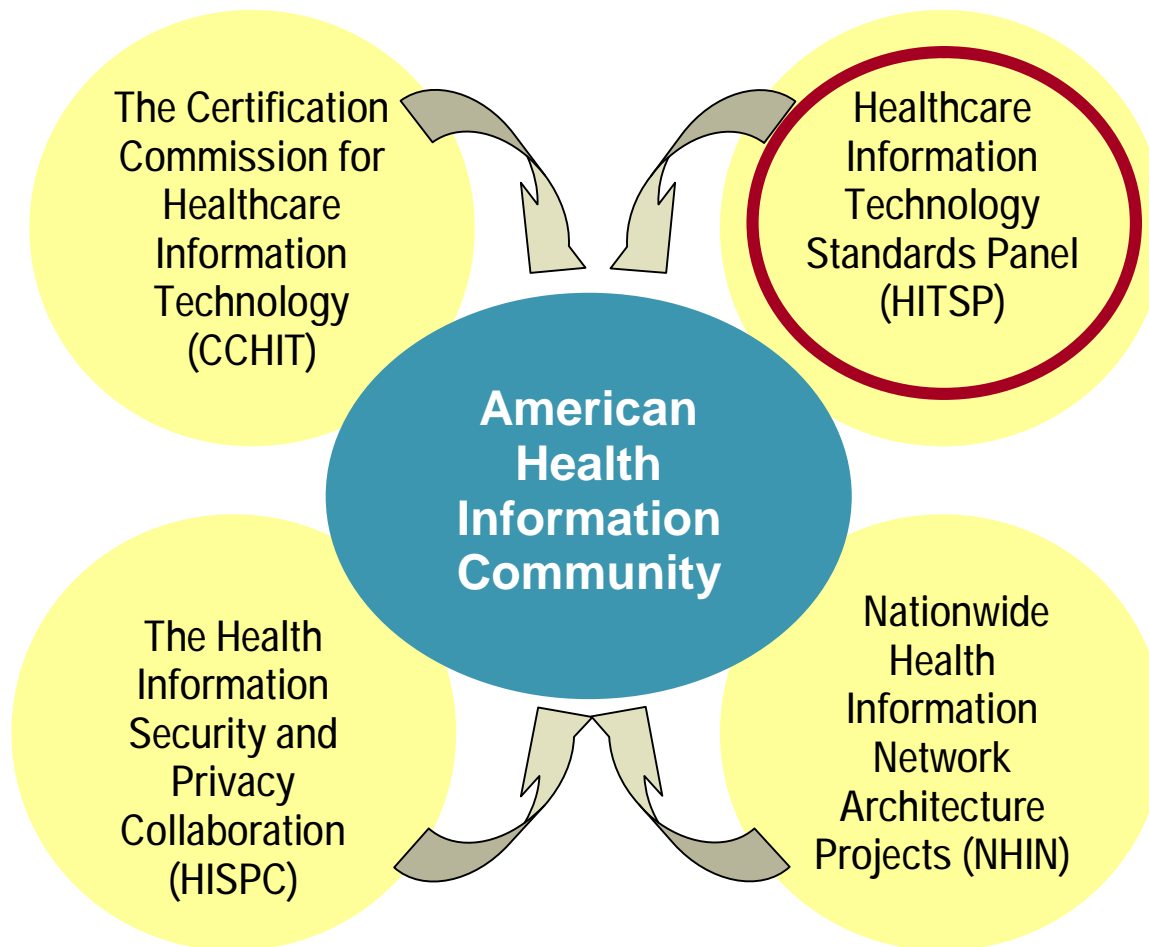
# Healthcare Information Technology Standards Panel

2006, 2007 and Beyond

John D. Halamka MD

Chair, HITSP

# A public-private “Community” was established to serve as the focal point for America’s health information concerns and drive opportunities for increasing interoperability



*HITSP includes 249 different member organizations and is administered by a Board of Directors*

- 16 SDOs (6%)
- 197 Non-SDOs (79%)
- 19 Govt. bodies (8%)
- 10 Consumer groups (4%)
- 7 Project Team and Undeclared (3%)

*The Community is a federally-chartered commission and will provide input and recommendations to HHS on how to make health records digital and interoperable, and assure that the privacy and security of those records are protected, in a smooth, market-led way.*



## 2006 – the First “Turn of the Crank”

### 4 Consumer Empowerment

- Medications
- Allergies
- Demographics
- Advance Directives

### 4 Electronic Health Records

- Laboratory including blood banking and microbiology
- Ordering and results exchange

### 4 Biosurveillance

- Deidentified registrations
- Labs
- Radiology text results

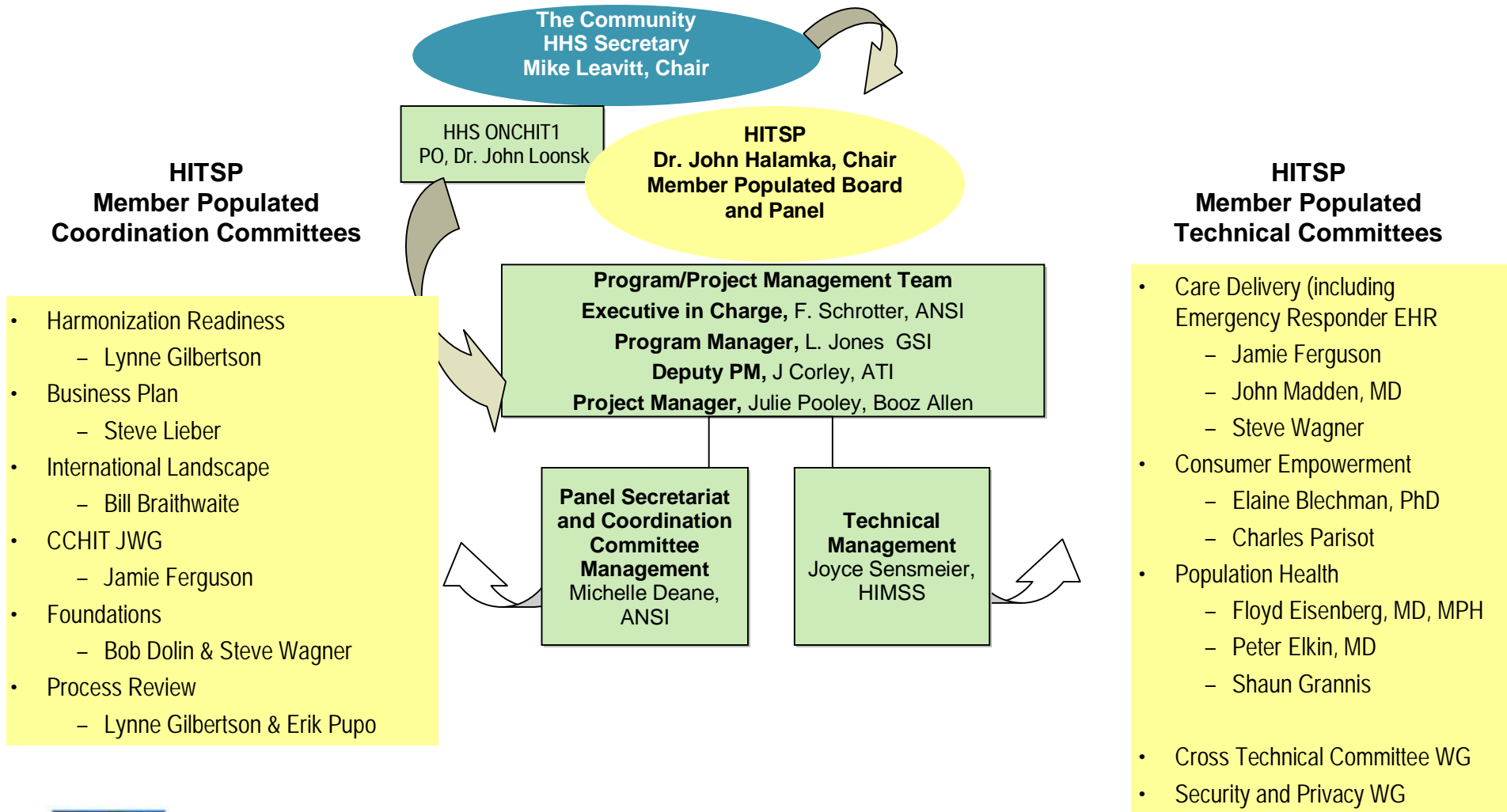


## What did we do in 2006?

- 4 Established the HITSP Organization and its committees
- 4 Created the standards harmonization process including all coordinating committee sub-processes
- 4 Harmonized standards for 3 use cases (accepted by Secretary Leavitt) and resolved three controversies along the way
  - To resolve CCR v. CDA, the CCD was successfully balloted
  - To resolve the need for Interim standards we accelerated CCD
  - ELINCS and an enhanced implementation guide for HL7 2.5.1 will be balloted by HL7 this Summer
- 4 To align HITSP interoperability specifications with CCHIT functional criteria, the CCHIT/HITSP Joint Working group is establishing a joint timeline for the next 3 years

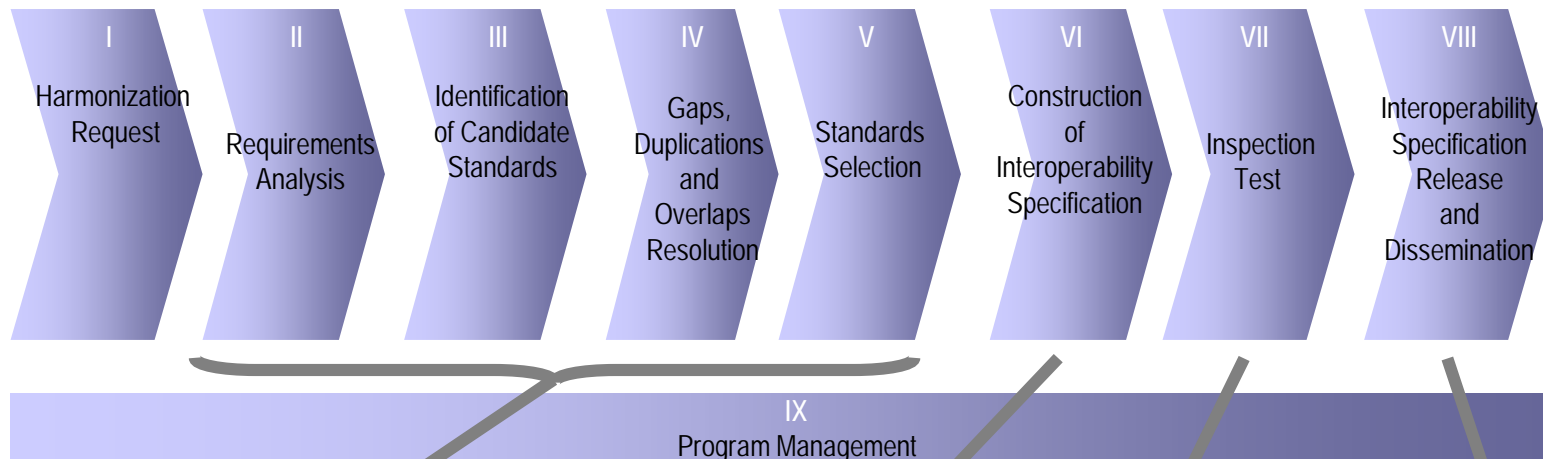


# HITSP Organization

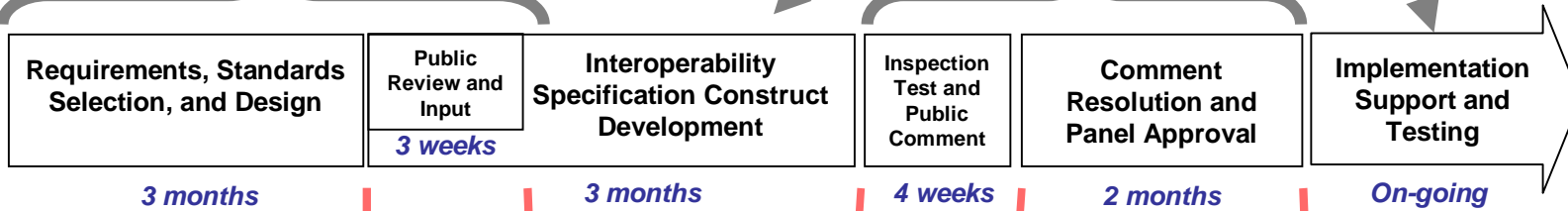


# Standards Harmonization Work Plan Tasks

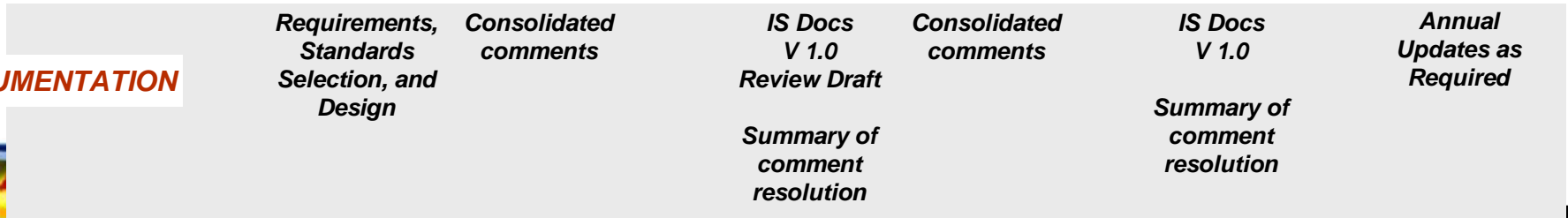
**PROCESS**



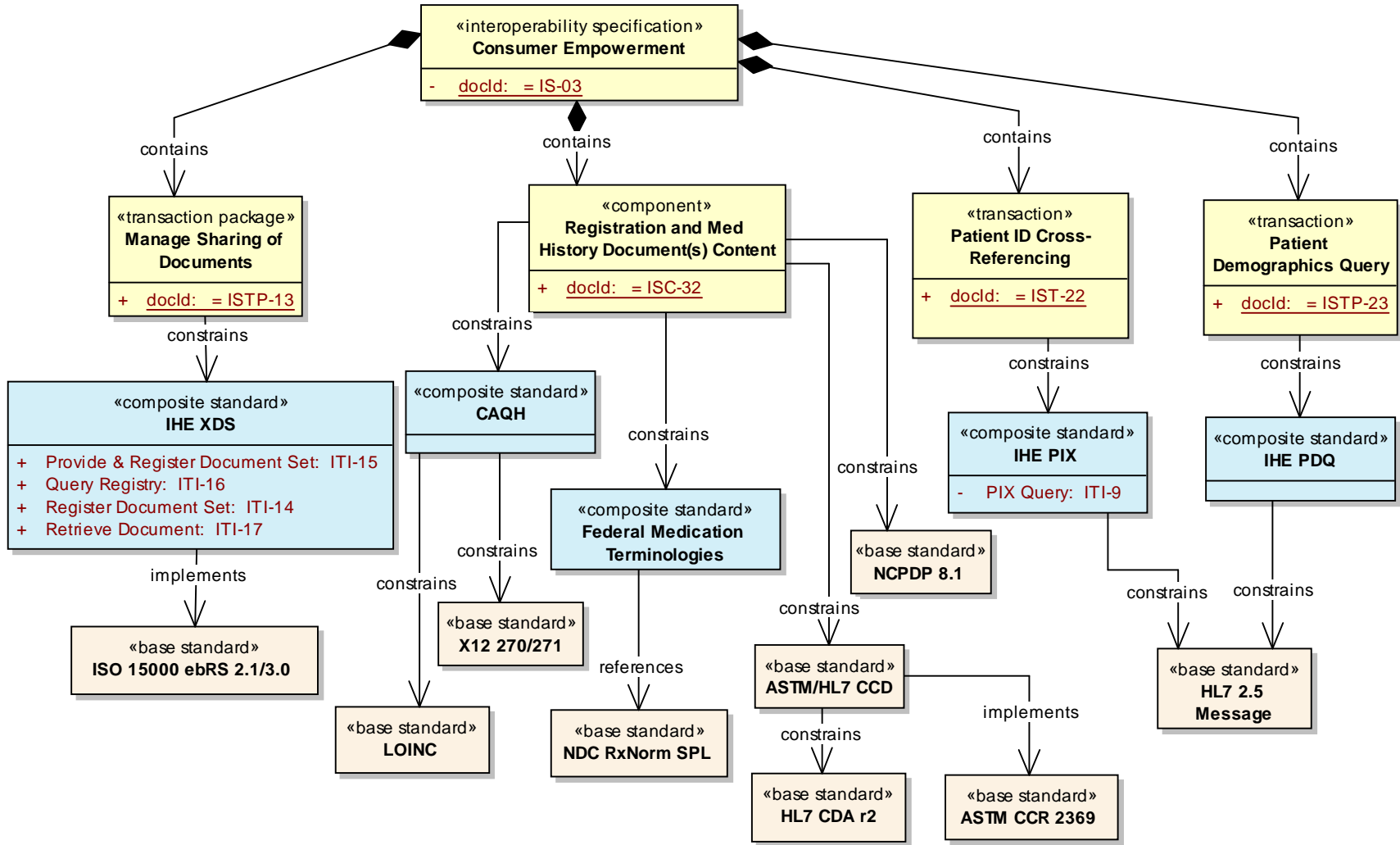
**TASKS**



**DOCUMENTATION**



cd CE Interoperability Specification



## ***HITSP Named Standards***

Accredited Standards Committee (ASC) X12 Insurance Subcommittee (X12N) Implementation Guides Version 004010 plus Addenda 004010A1

Accredited Standards Committee (ASC) X12 Standards Release 004010

American Society for Testing and Materials (ASTM) Standard Specification for Continuity of Care Record (CCR): # E2369-05

Council for Affordable Quality Health Care (CAQH) Committee on Operating Rules for Information Exchange (CORE) Phase I Operating Rules

Federal Medication Terminologies

Health Level Seven (HL7) Version 3.0 Continuity of Care Document (CCD)

Health Level Seven (HL7) Version 3.0 Clinical Document Architecture (CDA/CDA R2)

Health Level Seven (HL7) Version 2.5

Health Level Seven (HL7) EHR System Functional Model Draft Standard for Trial Use (DSTU)

Integrating the Healthcare Enterprise (IHE) Laboratory Technical Framework Supplement 2006-2007 Revision 1.0

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Integrating the Healthcare Enterprise (IHE) Patient Care Coordination (PCC) Technical Framework Revision 1.0

Logical Observation Identifiers Names and Codes (LOINC®)

National Council for Prescription Drug Programs (NCPDP) SCRIPT Standard Version 8.1



## 2007 – The second turn of the crank

- 4 Privacy and Security standards
- 4 Emergency Responder
- 4 Personal Health Records
- 4 Medication Management
- 4 Quality



# Security Standards

- 4 Collect and Communicate Security Audit Trails
- 4 Consistent Time
- 4 Document Integrity
- 4 Manage and Control Data Access
- 4 Manage and Control Privacy Consents
- 4 Manage Entity Identity Credentials
- 4 Non-Repudiation
- 4 Secure Communications Channel
- 4 User Authentication



# Security timeframe

- 4 First draft by June 18
- 4 Second draft and share those with other HISTP TCs for written comments June 20-July 8
- 4 Security and Privacy TC face to face meeting to finalize the documents July 10-11
- 4 Final documents to prepare for publication and public comment period by July 15
- 4 Inspection Test and Public Comment July 20 – August 16
- 4 Comment Resolution and Panel Approval August 17 – October 15



## Consumer Access to Clinical Information

|  |   |  |
|--|---|--|
| <b>C</b><br>Extension of existing Consumer Empowerment use case  | <p><b>Consumer Access to Clinical Information</b></p> <p>Consumers will benefit from the ability to access important healthcare data stored within their electronic health record to assist them in making decisions regarding care and healthy lifestyles. Accessible information could include registration information, medications history, lab results, current and previous health conditions, allergies, summaries of healthcare encounters and diagnoses. Consumers would be able to incorporate this information from their EHRs into Personal Health Records and share the information with designated individuals as needed. The PHR should describe medical terminology into layman's terms for the consumer. PHRs should be portable between vendors, so consumers can transfer the information as required.</p>   |  |
|  | <table style="width: 100%; border: none;"> <tr> <td style="vertical-align: top; width: 50%;"> <p><i>AHIC Priority Areas</i></p> <ul style="list-style-type: none"> <li>CE 1.0 Lab results as needed by patient</li> <li>CE 2.0 List of conditions and allergies</li> <li>CE 2.1 Health Problems</li> <li>CE 2.2 Medication Allergies</li> <li>CE 2.3 Allergies</li> <li>CE 6.2 Diagnosis codes</li> <li>AHIC 25.0 PHR portability methods</li> </ul> </td> <td style="vertical-align: top; width: 50%;"> <p><i>Workgroup Issues</i></p> <ul style="list-style-type: none"> <li>CE 11.0 Limited pre-populated clinical data &amp; limited patient access</li> <li>CE 13.0 Connectivity between physician offices, PHR's and pharmacies</li> <li>CE 12.0 Minimal interoperability or portability</li> <li>CE 10.0 PHR not integrated with workflow</li> <li>CE 14.0 State laws regarding labs</li> <li>CE 15.0 Policies for consumer entered data</li> </ul> </td> </tr> </table> | <p><i>AHIC Priority Areas</i></p> <ul style="list-style-type: none"> <li>CE 1.0 Lab results as needed by patient</li> <li>CE 2.0 List of conditions and allergies</li> <li>CE 2.1 Health Problems</li> <li>CE 2.2 Medication Allergies</li> <li>CE 2.3 Allergies</li> <li>CE 6.2 Diagnosis codes</li> <li>AHIC 25.0 PHR portability methods</li> </ul> |
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# Medications Management

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|--|--|
| <p style="text-align: center;">A</p> <p style="writing-mode: vertical-rl; transform: rotate(180deg);">New Use Case</p> | <p><b>Medications Management</b></p> <p>Consumers and providers would both benefit from electronic prescribing of medications, which would include transmittal of prescriptions to pharmacies by clinicians. Providers would be able to receive real-time feedback regarding potential adverse interactions and verify medication compliance by the consumer. Pharmacy Benefits Management entities would be able to interact with providers and consumers during the medications prescribing and fulfillment activities. Consumers would also be able to request prescription refills, view their prescription histories, verify insurance eligibility and coverage, view formulary information and incorporate all of this information into their Personal Health Records.</p> <p><i>AHIC Priority Areas</i></p> <ul style="list-style-type: none"> <li>CC 8.0 Monitoring of Medications</li> <li>EHR 2.0 Pharmacy/Allergy</li> <li>CCHIT 3.0 Medication management</li> <li>CCHIT 3.1 Outpatient prescription writing and transmission to pharmacies</li> <li>CCHIT 3.2 Ordering</li> <li>CCHIT 3.3 Clinical decision support</li> <li>CCHIT 3.4 Transmission</li> <li>CCHIT 3.5 Dispensing</li> <li>CCHIT 3.6 Administering</li> <li>CCHIT 3.7 Reconciliation</li> </ul> <p><i>Workgroup Issues</i></p> <ul style="list-style-type: none"> <li>EHR 11.0 Pharmacy/medication interoperability</li> <li>EHR 14.0 Need for confidentiality, privacy and security</li> </ul> |
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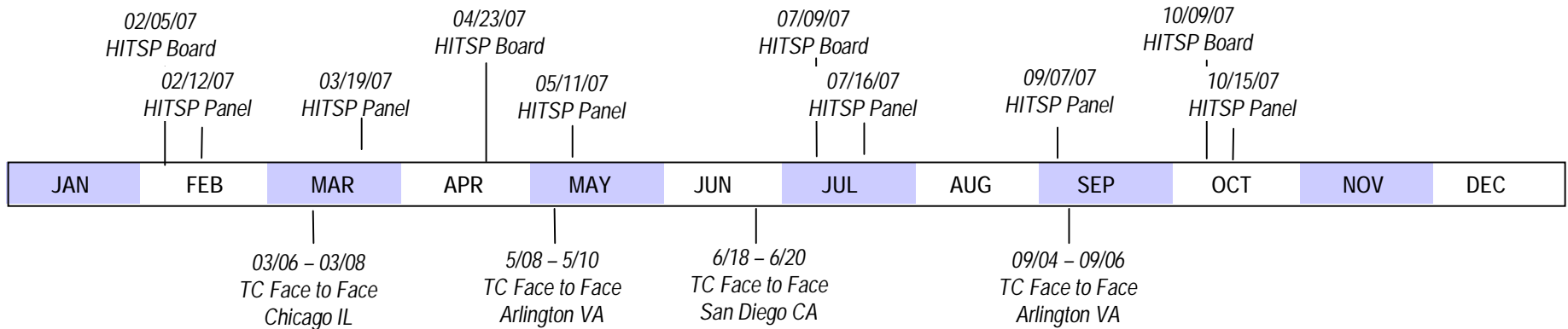


# Quality

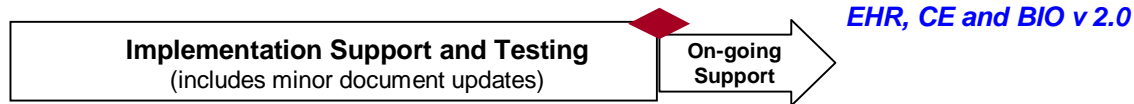
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|--|--|--|--|
| <p><b>A</b></p> <p>Extension of existing<br/>Biosurveillance use case</p>  | <p><b>Quality</b></p> <p>Providers would benefit from the collection and dissemination of healthcare quality data such as HQA quality indicators for inpatient care and AQA quality indicators for ambulatory care, particularly if this information can be integrated into EHR systems within the provider's workflows. Clinicians could benefit from receiving realtime or near-realtime feedback regarding relevant quality indicators and contra-indications for specific patients. Additionally, quality data across multiple providers and entities could be aggregated for the purpose of public reporting.</p> <table border="0"> <tr> <td data-bbox="453 862 1255 1081"> <p><i>AHIC Priority Areas</i></p> <p>Q 1.0 Inpatient Quality Measures (core set)</p> <p>Q 2.0 Ambulatory measures (core set)</p> <p>Q 3.0 Clinicians have access to feedback (self-assessment)</p> <p>Q 4.0 Public reporting</p> </td> <td data-bbox="1255 862 1902 1123"> <p><i>Workgroup Issues</i></p> <p>Q 8.0 Lack of data and technical standards and clinical documentation</p> <p>Q 9.0 Data sharing rights and responsibilities</p> <p>Q 10.0 Data security and privacy – policies for secondary use</p> </td> </tr> </table> | <p><i>AHIC Priority Areas</i></p> <p>Q 1.0 Inpatient Quality Measures (core set)</p> <p>Q 2.0 Ambulatory measures (core set)</p> <p>Q 3.0 Clinicians have access to feedback (self-assessment)</p> <p>Q 4.0 Public reporting</p> | <p><i>Workgroup Issues</i></p> <p>Q 8.0 Lack of data and technical standards and clinical documentation</p> <p>Q 9.0 Data sharing rights and responsibilities</p> <p>Q 10.0 Data security and privacy – policies for secondary use</p> |
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# HITSP 2007 Timeline



## Activity 1 – Version 2.0 of Existing EHR, CE, BIO ISs



## Activity 2 – On-going Work for Existing EHR, CE, BIO ISs (e.g. Security and Privacy) Activity 3 – New Emergency Responder EHR Use Case

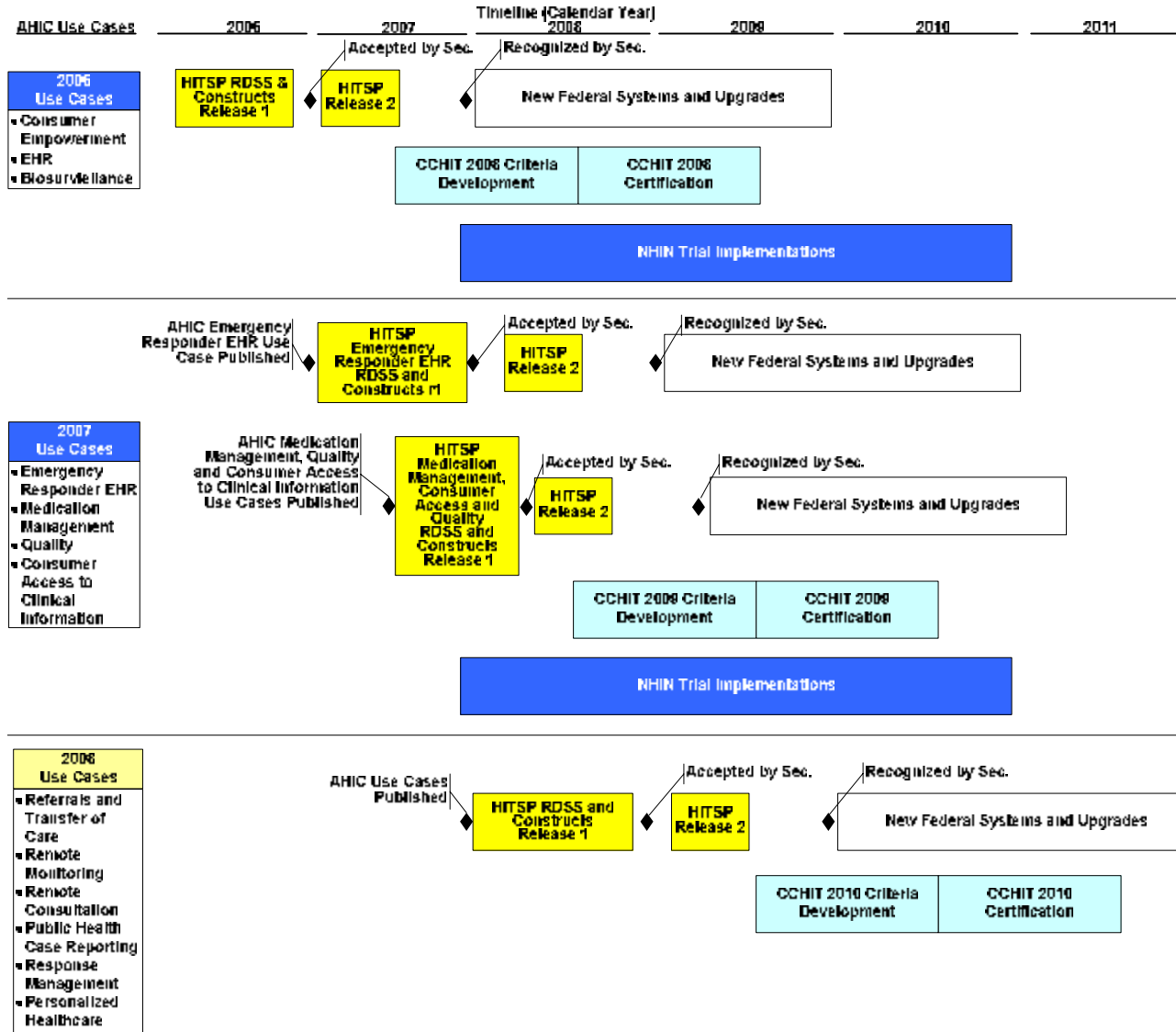
*S&P and EHR-ER v 1.0, EHR, CE, and BIO v M.m*



## Activity 4 –New Use Cases from AHIC



# Overall Timeline



## Summary

- 4 Over the past year, HITSP has become an established, trusted organization with a multi-stakeholder, open, transparent process for standards harmonization
- 4 In 2007, we will complete 4 additional use cases plus security/privacy
- 4 In 2008, we expect an additional 3-4 use cases
- 4 Our Foundations Committee will work on the medium to long term alignment of standards organizations and their work products in parallel with the Use Case work of the entire panel



## Contact Information

### 4 For general HITSP-related questions please contact:

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