







Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>									
						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
1	F	Identify and maintain a patient record: Key identifying information is stored and linked to the patient record. Both static and dynamic data elements will be maintained. A look up function uses this information to uniquely identify the patient.	1. The system shall create a single patient record for each patient.	DC.1.1.1	P				
2			2. The system shall associate (store and link) key identifier information (e.g., system ID, medical record number) with each patient record.	DC.1.1.1	P			Key identifier information must be unique to the patient record but may take any system defined internal or external form.	
3			3. The system shall provide the ability to store more than one identifier for each patient record.	DC.1.1.1	P			For interoperability, practices need to be able to store additional patient identifiers. Examples include an ID generated by an Enterprise Master Patient Index, a health plan or insurance subscriber ID, regional and/or national patient identifiers if/when such become available.	
4			4. The system shall use key identifying information to identify (look up) the unique patient record.	DC.1.1.1	P				
5			5. The system shall provide more than one means of identifying (looking up) a patient.	DC.1.1.1	P			Examples of identifiers for looking up a patient include date of birth, phone number.	
6			6. The system shall provide a field which will identify patients as being exempt from reporting functions.	DC.1.1.1	N			Examples include patients who are deceased, transferred, moved, seen as consults only. Being exempt from reporting is not the same as de-identifying a patient who will be included in reports. De-identifying patients for reporting is addressed in the "Health record output" functionality.	


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
8	F	Manage patient demographics: Contact information including addresses and phone numbers, as well as key demographic information such as date of birth, gender, and other information is stored and maintained for reporting purposes and for the provision of care.	1. The system shall capture and maintain demographic information as part of the patient record.	DC.1.1.2	P			Examples of a minimum set of demographic data elements include: name, address, phone number and date of birth. It is assumed that all demographic fields necessary to meet legislative and regulatory (e.g., HIPAA), research, and public health requirements will be included. A desirable feature would be a method of identifying how patients would like to be contacted (e.g., alternate addresses). De-identifying demographic information is addressed in the "Health record output" functionality.	
9			3. The system shall provide the ability to include demographic information in reports.	DC.1.1.2	P			This includes using demographics to generate reports and also allows demographics to be gathered into a report. See also "Report generation" functionality.	
10			The system shall provide the ability to maintain and make available historic information for demographic data including prior names, addresses, phone numbers and email addresses .	DC.1.1.2	N			Providers need this for look up and contact purposes, e.g., when attempting to locate a patient or family member for clinical communications.	
10a			The system shall maintain historic information for prior insurance.			N		Deleted	
11			5. The system shall provide the ability to modify demographic information about the patient.	DC.1.1.2	P				
12			6. The system shall store demographic information in the patient medical record in separate discrete data fields, such that data extraction tools can retrieve these data.	DC.1.1.2	N				
13	F		Manage problem list: Create and maintain patient specific problem	1. The system shall provide the ability to display all current problems associated with a patient.	DC.1.4.3	P			We assume current and active to mean the same thing.


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
14			lists.	2. The system shall provide the ability to maintain a history of all problems associated with a patient.	DC.1.4.3	P			This means both current and inactive and/or resolved problems. These may be viewed on separate screens or the same screen. Ideally each discrete problem would be listed once.
15				3. The system shall provide the ability to maintain the onset date of the problem.	DC.1.4.3	P			It is a vendor design decision whether to require complete date or free text of approximate date.
16				4. The system shall provide the ability to record the chronicity (chronic, acute/self-limiting, etc.) of a problem.	DC.1.4.3	P			
17				5. The system shall provide the ability to record the user ID and date of all updates to the problem list.	DC.1.4.3	P			
18				6. The system shall provide the ability to associate orders, medications, and notes with one or more problems.	DC.1.4.3	N			One should be able to identify all visits for a particular diagnosis/problem. Association can be made by free text notation or in structured data.
19				7. The system shall provide the ability to maintain a coded list of problems.	DC.1.4.3	P			For example, ICD-9, SNOMED-CT, DSM-IV. The Functionality WG will not specify which code set(s) are to be employed.
20				8. The system shall provide the ability to display inactive and/or resolved problems.		P			
21a				The system shall provide the ability to separately display active problems from inactive/resolved problems.		N			
22	F		Manage medication list: Create and maintain patient specific medication lists- Please see DC.1.7.1 for medication ordering as there is some overlap.	1. The system shall provide the ability to create and maintain medication lists.	DC.1.4.2	P			The medication list should be "patient-centric" and may include medications prescribed by any provider.
23				2. The system shall provide the ability to record the prescribing of medications including the identity of the prescriber.	DC.1.4.2	P			


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
24				3. The system shall provide the ability to maintain medication ordering dates.	DC.1.4.2	P			
25				4. The system shall provide the ability to maintain other dates associated with medications including start, modify, renewal and end dates as applicable.	DC.1.4.2	P			
26				5. The system shall provide the ability to display medication history for the patient.	DC.1.4.2	P			For clarification, medication history includes all medications prescribed since the EMR was established.
27				6. The system shall provide the ability to capture medications entered by authorized users other than the prescriber.	DC.1.4.2	P			It is important to have all current medications in the system for drug interaction checking. This in the future would include the incorporation of medication history obtained from outside electronic interfaces from insurers, PBMs, etc. "User" means medical and non-medical staff who are authorized by policy to enter prescriptions or other documentation.
28				7. The system shall provide the ability to enter non-prescription medications, including over the counter and complementary medications such as vitamins, herbs and supplements.	DC.1.4.2	P			This is important for interaction checking, associating symptoms with supplements e.g. the L-tryptophan related eosinophila-myalgia syndrome
29				8. The system shall provide the ability to exclude a medication from the current medication list (e.g., marked inactive, erroneous, completed, discontinued) and document reason for such action.	DC.1.4.2	P			Reason for removal or discontinuation may be captured as a discrete data element or as free text. In future this should be structured.
31				10. The system shall provide the ability to print a current medication list.	DC.1.4.2	P			
32				11. The system shall provide the ability to display current medications only.	DC.1.4.2	P			Excluding prior medications to make current medications easier to identify. Any given medication should display only once in the list.


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
33				12. The system shall include standard medication codes associated with each medication in the list. items in the medication list.	DC.1.4.2	N	N		It is anticipated that upcoming eRx regulation and the work of AHIC will define these in the near future. This requires publication by HITSP of an implementation guide by 3/07 3/06 . This requirement will be postponed for a year after the publication of such a guide if one is not available by 3/07 3/06 .
34				13. The system shall provide the ability to enter uncoded or free text medications when medications are not on the vendor-provided medication database –or information is insufficient to completely identify the medication.		M			Medications that are not on the vendor-provided medication database –or not enough information is available to completely identify the medication. This could be either uncoded (Synthroid unknown dose) or free text (blue hypertension pill)
35				14. The system shall provide the ability to alert the user at the time a new medication is prescribed that drug interaction and allergy checking will not be performed against the uncoded or free text medication.		N			
36				15. The system shall provide the ability to enter or further specify in a discrete field that the patient takes no medications.		N			
37				16. The system shall provide the ability to record the date of changes made to a patient's medication list and the identity of the user who made the changes. capture and display the identity of the user and date of changes made to the medication list for the patient.		M			This information may appear as an optional view rather than a required view on the main screen. Need to capture the identity of the user and the date of changes made.
38	F	Manage allergy and adverse reaction list: Create and maintain patient specific allergy and adverse		1.The system shall provide the ability to capture and store lists of medications and other agents to which the patient has had an allergic or other adverse reaction.	DC.1.4.1	P			The user determines what defines an allergy or adverse reaction.


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
39			reaction lists.	2. The system shall provide the ability to specify the type of allergic or adverse reaction.	DC.1.4.1	N			Allergy type may be specified as a discrete data element and/or as a free text description. This should be a modifiable field.
40				3. The system shall provide the ability to remove an item from the allergy and adverse reaction list.	DC.1.4.1	P			This could include removal, marking as erroneous, or marking as inactive. "Remove" in this context implies specifying that an allergy or allergen specification is no longer valid or active as opposed to deleting the information from the database entirely.
42a				5. The system shall provide the ability to record the removal of items from the allergy list. including the ID of the user who removed the item and attributes of the items removed.	DC.1.4.1	N			Necessary for medico-legal purposes. "Remove" in this context implies specifying that an allergy or allergen specification is no longer valid or active as opposed to deleting the information from the database entirely.
42b				The system shall provide the ability to record the identity of the user who added, modified, inactivated or removed items from the allergy list, including attributes of the changed items.		N			
43				6. The system shall provide the ability for a user to explicitly document that the allergy list was reviewed. The user ID and date stamp shall be recorded when the allergies reviewed option is selected. to review the allergies for a patient and record the date the review was performed and the ID of the user who performed it.	DC.1.4.1	M			Medico-legal and regulatory compliance
44				7. The system shall provide the ability to explicitly indicate that a patient has no known drug allergies.	DC.1.4.1	P			Medico-legal and regulatory compliance. This is meant to be specific to drug allergies.


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments	
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond		
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year				
46				9. The system shall provide the ability to capture non-drug agents to which the patient has had an allergic or other adverse reaction.	DC.1.4.1	P			These could include items such as foods or environmental agents. This need not be accomplished within the same portion of the chart where medication allergies are noted.	
47	F	Manage patient history: Capture, review, and manage medical, procedural/surgical, social and family history including the capture of pertinent positive and negative histories, patient reported or externally available patient clinical history.		1. The system shall provide the ability to capture, store, display, and manage patient history.	DC.1.2	P			Examples include past medical/surgical problems, diagnoses, procedures, family history and social history.	
48				2. The system shall provide the ability to capture structured data in the patient history.	DC.1.2	N			This function demonstrates the ability of a system to capture structured data but does not define the required elements of the patient history that shall be structured. Discrete data elements allow for searching and/or reporting by the EHR, and for this criterion the data could be free text or codified. Future functions would define the required patient history elements that shall be captured discretely as structured data, and where appropriate codified terminologies will be used.	
49					3. The system shall provide the ability to update a patient history by modifying, adding, removing, or inactivating items from the patient history as appropriate.	DC.1.2	P			Requirement not predicated on the capture of structured data.
50					4. The system shall provide the ability to capture patient history as both a presence and absence of conditions, i.e., the specification of the absence of a personal or family history of a specific diagnosis, procedure or health risk behavior.	DC.1.2	N			Requirement not predicated on the capture of structured data.


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
51				5. The system shall provide the ability to capture history collected from outside sources.	DC.1.2	P			This could include data from a personal health record, online patient histories, and information from pharmacy benefit management organizations. This criterion will accept any method of entry for year one, but electronic entry of information will be required thereafter.
53	F	Summarize health record		1. The system shall provide the ability to create and display a summary list for each patient that includes, at a minimum, the active problem list, current medication list, medication allergies and adverse reactions	DC.1.1.4	P			Health record summary is at the patient level as opposed to at the level of an individual visit or episode of care.
54	F	Manage clinical documents and notes: Create, correct, authenticate, and close, as needed, transcribed or directly entered clinical documentation.		1. The system shall provide the ability to create clinical documentation or notes (henceforth "documentation").	DC.1.9.1	P			
55				2. The system shall provide the ability to display documentation.	DC.1.9.1	P			
56				3. The system shall provide the ability to save a note in progress prior to finalizing the note.	DC.1.9.1	P			


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
57				4. The system shall provide the ability to finalize a note, i.e., change the status of the note from in progress to complete so that any subsequent changes are recorded as such.	DC.1.9.1	P			Medico-Legal. User rights are determined by role-based access defined in security. Only authorized users can complete, change or sign off a clinical note. The words, "sign," "signature," "cosign," and "cosignature" are intended here to convey actions, rather than referring to digital signature standards. It is recognized that an electronic signature is useful here. However, a widely accepted standard for electronic signatures does not exist. Thus, the criteria calls for documenting the actions of authenticated users at a minimum. In the future, when appropriate digital signature standards are available, certification criteria may be introduced using such standards.

 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology				Revisions from prev. release (18SEP06) are in red text			Discussion / Comments	
				Provisional Criteria (2007) are highlighted in yellow				
Compliance Key:			P = Previous Criteria					
			N = New for Year					
			M = Modified for Year					
Original line # Phase I	WG	Category and Description	Specific Criteria	Source or References	Compliance			
					Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
58			5. The system shall provide the ability to record the identity of the user finalizing each note and the date and time of finalization.	DC.1.9.1	P			Medico-Legal. User rights are determined by role-based access defined in security. Only authorized users can complete, change or sign off a clinical note. The words, "sign," "signature," "cosign," and "cosignature" are intended here to convey actions, rather than referring to digital signature standards. It is recognized that an electronic signature is useful here. However, a widely accepted standard for electronic signatures does not exist. Thus, the criteria calls for documenting the actions of authenticated users at a minimum. In the future, when appropriate digital signature standards are available, certification criteria may be introduced using such standards.
59			6. The system shall provide the ability to cosign a note and record the date and time of signature.		N			The words, "sign," "signature," "cosign," and "cosignature" are intended here to convey actions, rather than referring to digital signature standards. It is recognized that an electronic signature is useful here. However, a widely accepted standard for electronic signatures does not exist. Thus, the criteria calls for documenting the actions of authenticated users at a minimum. In the future, when appropriate digital signature standards are available, certification criteria may be introduced using such standards.

Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
60				7. The system shall provide the ability to addend and/or correct notes that have been finalized.	DC.1.9.1	P			The words, "sign," "signature," "cosign," and "cosignature" are intended here to convey actions, rather than referring to digital signature standards. It is recognized that an electronic signature is useful here. However, a widely accepted standard for electronic signatures does not exist. Thus, the criteria calls for documenting the actions of authenticated users at a minimum. In the future, when appropriate digital signature standards are available, certification criteria may be introduced using such standards.
61				8. The system shall provide the ability to record and display the identity of the user who addended or corrected a note, as well as other attributes of the addenda or correction, such as the date and time of the change.	DC.1.9.1	P			Necessary for medico-legal purposes. The words, "sign," "signature," "cosign," and "cosignature" are intended here to convey actions, rather than referring to digital signature standards. It is recognized that an electronic signature is useful here. However, a widely accepted standard for electronic signatures does not exist. Thus, the criteria calls for documenting the actions of authenticated users at a minimum. In the future, when appropriate digital signature standards are available, certification criteria may be introduced using such standards.
62				9. The system shall provide the ability to enter free text notes.	DC.1.9.1	P			

Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
63				10. The system shall provide the ability to filter, search or order notes by the provider who finalized the note.	DC.1.9.1	N			
64				11. The system shall provide the ability to filter, search or order notes by associated diagnosis within a patient record.	DC.1.9.1	N			This is intended to be the coded diagnosis and not free text in the body of a note.
65				12. The system shall provide the ability to capture patient vital signs, including blood pressure, heart rate, respiratory rate, height, and weight, as discrete data.	DC.1.9.1	P			It is understood that vendors should support conversion to numeric values that can be graphed. Coding in ICD9/10, SNOMED, UMLS, etc., would enhance interoperability and for public health surveillance or clinical research.
68				15. The system shall provide templates for inputting data in a structured format as part of clinical documentation.	DC.1.9.1	P			Codified data are data that is structured AND codified according to some 'external' industry accepted standard such as ICD-9, SNOMED-CT, and CPT-4.
69				16. The system shall provide the ability to customize clinical templates.	DC.1.9.1	P			Customizations may be site specific.
71a				The system shall be capable of recording comments by the patient or the patient's representative regarding the accuracy or veracity of information in the patient record (henceforth 'patient annotations').		N			For 2007 it is sufficient for these to be recorded as either free-text notes (see item F54) or scanned paper documents (see item F78). It is not required that the system facilitate direct entry into the system by the patient or patient's representative.
74				21. The system shall provide the ability to graph height and weight over time.		P			

Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
76	F	Capture external clinical documents: Incorporate clinical documentation from external sources.	1. The system shall provide the ability to capture and store external documents.	DC.1.1.3.1	P			Scanned documents are sufficient in 2005, granular data will be expected in the future. This covers all types of documents received by the practice that would typically be incorporated into a medical record, including but not limited to faxes, referral authorizations, consultant reports, and patient correspondence of a clinical nature.	
77			2. The system shall provide the ability to receive, store in the patient's record, and display discrete lab results received through an electronic interface.	DC.1.1.3.1	P			This may be an external source such as a commercial lab or through an interface with on site lab equipment.	
78			3. The system shall provide the ability to save scanned documents as images.	DC.1.1.3.1	P				
79			4. The system shall provide the ability to receive, store in the patient's record, and display text-based outside reports.	DC.1.1.3.1	P			This could be either from an outside system or from scanning with optical character recognition. Integration here means the ability to find and display the documents within the system.	
85	F	Generate and record patient specific instructions: Generate and record patient specific instructions as clinically indicated.	1. The system shall provide access to patient instructions and patient educational materials, which may reside within the system or be provided through links to external sources.	DC.1.10	N			An example would be a vaccine information statement.	
86			2. The system shall have the ability to provide access to medication instructions, which may reside within the system or be provided through links to external sources.	DC.1.10	P				


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
87				3. The system shall have the ability to provide access to test and procedure instructions that can be customized by the physician or health organization. These documents may reside within the system or be provided through links to external sources.	DC.1.10	M			This item relates to customization of instructions, not to recording in patient record that instructions have been provided.
88				4. The system shall provide the ability to record that patient specific instructions or educational material were provided to the patient.	DC.1.10	P			This does not require automatic documentation.
89				5. The system shall provide the ability to create patient specific instructions.	DC.1.10	P			
90	F	Order medication: Create prescriptions or other medication orders with detail adequate for correct filling and administration.		1. The system shall provide the ability to create prescription or other medication orders with sufficient information for correct filling and administration by a pharmacy.	DC.1.7.1	P			The term pharmacy here refers to all entities which fill prescriptions and dispense medications including but not limited to retail pharmacies, specialty, and mail order pharmacies.
92				3. The system shall provide the ability to record user and date stamp for prescription related events, such as initial creation, renewal, refills, discontinuation, and cancellation of a prescription.	DC.1.7.1	P			Security to limit prescription writing is included in I.1.2 below.



AMBULATORY FUNCTIONALITY
Proposed 2007 Final Criteria - Feb 14 2007
 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006
 © 2007 The Certification Commission for Healthcare Information Technology

Revisions from prev. release (18SEP06) are in red text
 Provisional Criteria (2007) are highlighted in yellow
 Compliance Key:
 P = Previous Criteria
 N = New for Year
 M = Modified for Year

Original line # Phase I	WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
					Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
93			4. The system shall provide the ability to capture the identity of the prescribing provider for all medication orders	DC.1.7.1	P			The words, "sign," "signature," "cosign," and "cosignature" are intended here to convey actions, rather than referring to digital signature standards. It is recognized that an electronic signature is useful here. However, a widely accepted standard for electronic signatures does not exist. Thus, the criteria calls for documenting the actions of authenticated users at a minimum. In the future, when appropriate digital signature standards are available, certification criteria may be introduced using such standards.
95			6. The system shall provide the ability to update the medication history with the newly prescribed medications.	DC.1.7.1	P			The words, "sign," "signature," "cosign," and "cosignature" are intended here to convey actions, rather than referring to digital signature standards. It is recognized that an electronic signature is useful here. However, a widely accepted standard for electronic signatures does not exist. Thus, the criteria calls for documenting the actions of authenticated users at a minimum. In the future, when appropriate digital signature standards are available, certification criteria may be introduced using such standards.
96			7. The system shall have the ability to provide a list of medications to search from, including both generic and brand name.	DC.1.7.1	N			


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
97				8. The system shall provide the ability to maintain a coded list of medications.	DC.1.7.1	P			For clarification - Coding means a unique identifier for each medication. This functional requirement does not intend to require a national system of coding for medications.
98				9. The system shall provide the ability to capture common content for prescription details including strength, sig, quantity, and refills to be selected by the ordering clinician.	DC.1.7.1	P			We encourage the development of standard national abbreviations and that only approved abbreviations should be supported.
103				14. The system shall provide the ability to reorder a prior prescription without re-entering previous data (e.g. administration schedule, quantity).	DC.1.7.1	P			
104				15. The system shall provide the ability to print and electronically fax prescriptions.	DC.1.7.1	P			Appropriate audits and security should be in place.
105				16. The system shall provide the ability to re-print and re-fax prescriptions.		P			This allows a prescription that did not come out of the printer, or a fax that did not go through, to be resent/reprinted without entering another prescription. Appropriate audits and security should be in place.
106				17. The system shall provide the ability to submit prescriptions electronically.	DC.1.7.1	N			See also line 166 (DC 3.2.2). Faxing for 2006, tentative electronic 2007 once standards are promulgated. This presupposes that the pharmacy is capable of receiving electronic prescriptions. This function relates to computer e-prescribing and not faxing. Appropriate audits and security should be in place.
110				21. The system shall provide the ability to identify medication samples dispensed, including lot number and expiration date.	DC.1.7.1	N			Lot numbers and expiration date could be entered in free text or encoded.





AMBULATORY FUNCTIONALITY
Proposed 2007 Final Criteria - Feb 14 2007
Proposed FUNCTIONALITY For 2007 Certification of
Ambulatory EHRs incorporates AFWG work to 28 Nov 2006
 © 2007 The Certification Commission for Healthcare Information Technology


Revisions from prev. release (18SEP06) are in red text
 Provisional Criteria (2007) are highlighted in yellow
 Compliance Key:
 P = Previous Criteria
 N = New for Year
 M = Modified for Year


Original line # Phase I	WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
					Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
111			22. The system shall provide the ability to prescribe fractional amounts of medication (e.g. 1/2 tsp, 1/2 tablet).	DC.1.7.1	P			Very important to prescribing for pediatric and geriatric patients.
112			23. The system shall provide the ability to prescribe uncoded medications.		N			See DC.1.4.2
113			24. The system shall provide the ability to alert the user at the time a new medication is prescribed that drug interaction, allergy, and formulary checking will not be performed against the uncoded medication.		N			
114			25. The system shall provide the ability to update drug interaction databases.		P			This includes updating or replacing the database with a current version.
115			26. The system shall provide the ability to alert the user if the drug interaction information is outdated.		M	N		The drug database should have an "expiration date" based on the frequency of their updates such that when that date has passed, the user is alerted.
116			27. System shall provide the ability to allow the user to configure prescriptions to incorporate fixed text according to the user's specifications and to customize the printed output of the prescription.		P			This refers to the "written" output and language on the prescription such as specific language, dispense as written. For instance, users should be able to modify the format/content of printed prescriptions to comply with state Board of Pharmacy requirements.
117			28. The system shall provide the ability to associate a diagnosis with a prescription.		P			
118			29. The system shall provide the ability to display the associated problem or diagnosis (indication) on the printed prescription.		M			At least one diagnosis shall be able to be displayed but the ability to display more than one is desirable. Associated problem or diagnosis can be free text or structured data.


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
120				31. The system shall provide the ability to create provider specific medication lists of the most commonly prescribed drugs with a default dose, frequency, and quantity.		N			
122	F	Order diagnostic tests: Submit diagnostic test orders based on input from specific care providers.		1. The system shall provide the ability to order diagnostic tests, including labs and imaging studies.	DC.1.7.2.2	P			This includes physicians and authorized non-physicians.
124				3. The system shall provide the ability to capture the identity of the ordering provider for all test orders.		P			
126				5. The system shall provide the ability to capture appropriate order entry detail, including associated diagnosis.	DC.1.7.2.2	P			Including associated diagnoses. It is desirable that all information for medical necessity checking be captured.
127				6. The system shall provide the ability to display instructions and/or prompts created by the user when ordering diagnostic tests or procedures. to the ordering user when placing orders for diagnostic tests so that the user supplies all required information.	DC.1.7.2.2	N			Does not require vendor to set required fields, but it is a highly desirable function.
128				7. The system shall provide the ability to relay orders for a diagnostic test to the correct destination for completion.	DC.1.7.2.2	P			Mechanisms for relaying orders may include providing a view of the order, sending it electronically, or printing a copy of the order or order requisition.
129				8. The system shall have the ability to provide a view of active orders for an individual patient.	DC.1.7.2.2	N			Additional sorts and filters may be provided by the vendors but not required.
130				9. The system shall have the ability to provide a view of orders by like or comparable type, e.g., all radiology or all lab orders.	DC.1.7.2.2	N			May include filters or sorts.
131	F			Manage order sets: Provide order sets based on provider input or system prompt, medication suggestions, drug recall updates.		1. The system shall provide the ability to define a set of related orders to be subsequently ordered as a group on multiple occasions.	DC.1.7.3	N	
132		2. The system shall provide the ability to modify order sets.	DC.1.7.3			N			


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
133				3. The system shall provide the ability to include in an order set orders for medications, laboratory tests, imaging studies, procedures and referrals.	DC.1.7.3	N			
134				4. The system shall provide the ability to display orders placed through an order set either individually or as a group.	DC.1.7.3	N			Need to be able to see the individual components of the order set, rather than just the name of the order set. Does not mean to break down a lab panel into individual components.
136	F	Manage results: Route, manage, and present current and historical test results to appropriate clinical personnel for review, with the ability to filter and compare results.		1. The system shall provide the ability to indicate normal and abnormal results based on data provided from the original data source.	DC.1.8.3	P			As each lab has it's own normal values, these should be reflected in the indication as to whether a lab is normal or abnormal.
137				2. The system shall provide the ability to display numerical results in flow sheets and graphical form in order to compare results, and shall provide the ability to display values graphed over time.	DC.1.8.3	N			It is desirable for the system indicate if abnormal results are high or low.
138				3. The system shall provide the ability to display non-numeric current and historical test results as textual data.	DC.1.8.3	P			
139				4. The system shall provide the ability to notify the relevant providers (ordering, copy to) that new results have been received.	DC.1.8.3	N			Examples of notifying the provider include but are not limited to a reference to the new result in a provider "to do" list or inbox.
140a				The system shall provide the ability to filter or sort results by type of test and test date.		N			
140b				In areas where results from multiple patients are displayed, the system shall provide the ability to filter or sort results by patient.		N	N		
141				6. The system shall provide the ability to forward a result to other users.	DC.1.8.3	N			
142				7. The system shall provide the ability to transfer the responsibility to perform follow up actions from clinical to other clinical personnel.	DC.1.8.3	N			Deleted.


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>									
						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
143				8. The system shall provide the ability to link the results to the original order.	DC.1.8.3	N			In 2007 this link can be effected manually by changing the status of the order from pending to complete. Future requirements could automate this link for certain electronically received labs although the requirement should not require that all types of orders be electronically linked to the results since the variety of result formats can be quite large (PT consult, Diabetes education...) and even the variety of lab result formats can be wide.
144				9. The system shall provide the ability for a user to attach a free text comment to a result that can be seen by another user who might subsequently view that result. to enter a free text annotation to a result.	DC.1.8.3	N			
146				11. The system shall provide the ability for a user to whom a result is presented to acknowledge the result.	DC.1.8.3	P			This is separate from audit trail.
147	F	Manage consents and authorizations: Create, maintain, and verify patient treatment decisions in the form of consents and authorizations when required.		1. The system shall provide the ability to capture scanned paper consent documents (covered in DC.1.1.3.1).	DC.1.3.3	P			
148				2. The system shall provide the ability to store, display and print patient consent forms. generate both on-line on-screen and printable consent forms.	DC.1.3.3	M			Example: Consent forms stored in the computer which are capable of being signed by the patient with either an electronic pen or a digital signature once widely available.
149					3. The system shall provide the ability to store and display administrative authorizations (e.g. privacy notices).	DC.1.3.3	N		


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
152	F	Manage patient advance directives: Capture, maintain, and provide access to patient advance directives.	1. The system shall provide the ability to indicate that a patient has completed advanced directive(s).	DC.1.3.2	P			Important for appropriate use of resources at end of life and may just include a yes, no indication.	
153			2. The system shall provide the ability to indicate the type of advanced directives, such as living will, durable power of attorney, or a "Do Not Resuscitate" order.	DC.1.3.2	N			This may be recorded in free text or as discrete data.	
154			3. The system shall provide the ability to indicate when advanced directives were last reviewed.	DC.1.3.2	N			This may be recorded in free text or as discrete data.	
155	F	Support for standard care plans, guidelines, protocols: Support the use of appropriate standard care plans, guidelines, and/or protocols for the management of specific conditions.	1. The system shall have the ability to provide access to standard care plan, protocol and guideline documents when requested at the time of the clinical encounter. These documents may reside within the system or be provided through links to external sources.	DC.2.2.1.1	P			This requirement could be met by simply including links or access to a text document. Road map would require more comprehensive decision support in the future. This includes the use of clinical trial protocols to ensure compliance.	
156			2. The system shall provide the ability to create site-specific care plan, protocol, and guideline documents.	DC.2.2.1.1	P			This includes the use of clinical trial protocols to ensure compliance. It is expected that in the future discrete data elements from other areas of the chart will populate matching fields.	
157			3. The system shall provide the ability to modify site-specific standard care plan, protocol, and guideline documents obtained from outside sources.	DC.2.2.1.1	N				
159	F	Capture variances from standard care plans, guidelines, protocols: Identify variances from patient-	1. The system shall provide the ability to record the reason for variation from care plans, guidelines, and protocols as discrete data.		N		2009	Needed for pay for performance.	


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
160	F	Support for drug interaction: Identify drug interaction warnings at the point of medication ordering	1. The system shall provide the ability to check for potential interactions between medications to be prescribed and current medications and alert the user at the time of medication ordering if potential interactions exist.	DC.2.3.1.1	P			This reduces risk of inappropriate prescribing, prevents pharmacy call backs, and can reduce malpractice liability.	
161			2. The system shall provide the ability to check for potential interactions between medications to be prescribed and medication allergies and intolerances listed in the record and alert the user at the time of medication ordering if potential interactions exist.	DC.2.3.1.1	P				
162			3. The system shall provide the ability to prescribe a medication despite alerts for interactions and/or allergies being present.	DC.2.3.1.1	P				
163			4. The system shall provide the ability to set the severity level at which drug interaction warnings should be displayed.	DC.2.3.1.1	P				
165			The system shall provide the ability to document at least one reason reasons for overriding a drug any drug drug or drug-allergy interaction warning triggered at the time of medication ordering.	DC.2.3.1.1	N			Necessary for medico-legal purposes.	
168			9. The system shall provide the ability to display, on demand, potential interactions on a patient's medication list, even if a medication is not being prescribed at the time.	DC.2.3.1.1	N				
173	F		Support for medication or immunization administration or supply: To reduce medication errors at the time of administration of a medication, the patient is positively	1. The system shall provide the ability to document medication administration.	DC.2.3.2	P			
174		2. The system shall provide the ability to document immunization administration.		DC.2.3.2	P				


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>									
						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
175			identified; checks on the drug, the dose, the route and the time are facilitated. Documentation is a by-product of this checking; administration details and additional	3. The system shall provide the ability to document, for any immunization, the immunization type, dose, time of administration, route, site, lot number, expiration date, manufacturer, and user ID as structured documentation.	DC.2.3.2	N			Capturing this information for non-immunizations is optional.
178	F	Support for non-medication ordering (referrals, care management)		1. The system shall provide the ability to create referral orders with detail adequate for correct routing.	DC.2.4.2	N			This could include referrals to sub-specialists, physical therapy, speech therapy, nutritionists, and other non-medication, non-clinical order. Adequate detail includes but is not limited to: • Date • Patient name and identifier • "Refer to" specialist name, address and telephone number • "Refer to" specialty • Reason for referral • Referring physician name
179				2. The system shall provide the ability to record user ID and date/time stamp for all referral related events.	DC.2.4.2	N			Necessary for medico-legal purposes.
180	F	Present alerts for disease management, preventive services and wellness: At the point of clinical decision making, identify		1. The system shall provide the ability to establish criteria for disease management, wellness, and preventive services based on patient demographic data (minimally age and gender).	DC.2.5.1	P			This includes the use of clinical trial protocols to ensure compliance.


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>									
<p>Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year</p>									
181			patient specific suggestions/reminders, screening tests/exams, and other preventive services in support of disease management, routine preventive and wellness patient care standards.	2. The system shall provide the ability to display alerts based on established guidelines.	DC.2.5.1	P			Guidelines may be from national organizations, payers, or internal protocols. It is expected that in the future discrete data elements from other areas of the chart will populate matching fields. It is assumed that when a service is completed, this change will be immediately reflected with removal of the prompt.
182				3. The system shall provide the ability to establish criteria for disease management, wellness, and preventive services based on clinical data (problem list, current medications).	DC.2.5.1	P			Lab results in future years
183				4. The system shall provide the ability to update disease management guidelines and associated reference material.	DC.2.5.1	N			This allows the system's decision support tools to support changes in best practice guidelines.
184				5. The system shall provide the ability to update preventive services/wellness guidelines and associated reference material.	DC.2.5.1	P			
185				6. The system shall provide the ability to override guidelines.	DC.2.5.1	P			
186				7. The system shall provide the ability to document reasons disease management or preventive services/wellness prompts were overridden.	DC.2.5.1	N			Needed for some pay for performance initiatives. Needed for medico-legal reasons and clinical decision support.
187				8. The system shall provide the ability to modify the rules or parameters upon which guideline-related alerts are based. guidelines.	DC.2.5.1	N			This is necessary for modifications as guidelines change or practices wish to adhere to more stringent levels for example, using a HbA1c target of 6.5% instead of 7%.


Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
188				9. The system shall provide the ability to document that a preventive or disease management service has been performed based on activities documented in the record (e.g., vitals signs taken).	DC.2.5.1	N			
189				10. The system shall provide the ability to document that a disease management or preventive service has been performed with associated dates or other relevant details recorded.	DC.2.5.1	N			This could include services performed internally or external to the practice.
189a				11. The system shall provide the ability to individualize alerts to address a patient's specific clinical situation.		M			This is done at the patient level. Examples include but are not limited to: • Remove mammography for woman that has had a mastectomy • Remove annual pap smear alert for a woman who has had a complete hysterectomy. • Inactivate an alert for routine colon cancer screening in a patient who is terminally ill.
190	F	Notifications and reminders for disease management, preventive services and wellness: Between healthcare encounters, notify the patient and/or appropriate provider of those preventive services, tests, or behavioral actions that are due or overdue.		1. The system shall provide the ability to identify preventive services, tests, or counseling that are due on an individual patient.	DC.2.5.2	P			In the future, the system should perform this automatically and proactively "contact" patient(s) without physician intervention (e.g. automated reminder letter). These guidelines might come from national organizations, medical societies, etc.
191				2. The system shall provide the ability to display reminders for disease management, preventive, and wellness services in the patient record.	DC.2.5.2	P			It is expected that in the future discrete data elements from other areas of the chart will populate matching fields.
192				3. The system shall provide the ability to identify criteria for disease management, preventive, and wellness services based on patient demographic data (age, gender).	DC.2.5.2	P			

Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>									
						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
193				4. The system shall provide the ability to identify criteria for disease management, preventive, and wellness services based on clinical data (problem list, current medications, lab values).	DC.2.5.2	N			These guidelines could be tailored to address payer-specific criteria but we would encourage national standardization of guidelines.
194				5. The system shall provide the ability to modify the guidelines that trigger the reminders.	DC.2.5.2	P			
195				6. The system shall provide the ability to notify the provider that patients are due or are overdue for disease management, preventive, or wellness services.	DC.2.5.2	P			
196				7. The system shall provide the ability to produce a list of patients who are due or are overdue for disease management, preventive, or wellness services.	DC.2.5.2	P			
198	F	Clinical task assignment and routing: Assignment, delegation and/or transmission of tasks to the appropriate parties.		1. The system shall provide the ability to create and assign tasks by user or user role.	DC.3.1.1	P			Examples of tasks are messages, notifications, inbox items, worklist to-do's. This task assignment refers to internal users. External tasks would be handled under ordering section.
199				2. The system shall provide the ability to present a list of tasks by user or user role.	DC.3.1.1	N			Examples of tasks are messages, notifications, inbox items, worklist to-do's. This task assignment refers to internal users. External tasks would be handled under ordering section.
200				3. The system shall provide the ability to re-assign and route tasks from one user to another user.	DC.3.1.1	N			
201				4. The system shall provide the ability to designate a task as completed.	DC.3.1.1	P			
202				5. The system shall provide the ability to remove a task without completing the task.	DC.3.1.1	P			Removing a task eliminates it from an individual user's "to do" list, not from audit logs, etc.

Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
204	F	Inter-provider communication: Support secure electronic communication (inbound and outbound) between providers in the same practice to trigger or respond to pertinent actions in the care process (including referral), document non-electronic communication (such as phone calls, correspondence or other encounters) and generate paper message artifacts where appropriate.	1. The system shall provide the ability to document verbal/telephone communication into the patient record.	DC.3.2.1	P				
205			2. The system shall provide the ability to incorporate paper documents from external providers into the patient record.	DC.3.2.1	P				
206			3. The system shall support messaging between users.	DC.3.2.1	P			Results and other patient data could be included. As clarification, messaging is defined as any text string sent from one person to another in the office.	
207	F	Pharmacy communication: Provide features to enable secure and reliable communication of information electronically between practitioners and pharmacies or between practitioner and intended recipient of pharmacy orders.	1. The system shall have the ability to provide electronic communication between prescribers and pharmacies or other intended recipients of the medication order.	DC.3.2.2	P			Until electronic standards are established, FAX is a suitable means of transmission.	
208			2. The system shall provide the ability to electronically communicate from the prescriber to the pharmacy an initial medication order as well as changes to or renewals of an existing order.	DC.3.2.2	N			Cancellations would be included in this function.	
209			3. The system shall capture any acknowledgments, prior authorizations, renewals inquiries and fill notifications provided by the pharmacy or other participants in the electronic prescription. The system shall provide the ability to capture and display any renewal requests received electronically from or on behalf of any dispensing entity.	DC.3.2.2	N	N			

Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
210	F	Provider demographics: Provide a current directory of practitioners that, in addition to demographic information, contains data needed to determine levels of access required by the EHR security and to support the practice of medicine.	1. The system shall provide the ability to maintain a directory of all clinical personnel who currently use or access the system.	S.1.3.1	P				
211			2. The system shall provide the ability to maintain a directory which contains identifiers required for licensed clinicians to support the practice of medicine including a minimum state medical license, DEA, NPI, and UPIN number.	S.1.3.1	N			This directory may be the same as that in criteria #1 for this functionality.	
212			3. The system shall provide the ability to maintain a directory that stores user attributes required to determine the system security level to be granted to each user.	S.1.3.1	P			This directory may be the same as that in criteria #1 for this functionality.	
213			4. The system shall allow authorized users to update the directory.	S.1.3.1	P				
214			5. The system shall provide the ability to create and maintain a directory of clinical personnel external to the organization who are not users of the system to facilitate communication and information exchange.	S.1.3.1	M			This directory may be the same as that in criteria #1 for this functionality.	
215	F	Scheduling: Support interactions with other systems, applications, and modules to provide the necessary data to a scheduling system for optimal efficiency in the scheduling of patient care, for either the patient or a resource/device.	1. The system shall provide the ability to display a schedule of patient appointments, populated either through data entry in the system itself or through an external application interoperating with the system.	S.1.6	P				
216	F	Report Generation: Provide report generation features for the generation of standard and ad hoc reports	1. The system shall provide the ability to generate reports of clinical and administrative data using either internal or external reporting tools.	S.2.2	N			Needed for pay for performance, quality improvement activities. All data that is entered in a structured format should be individually reportable.	

Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
217				2. The system shall provide the ability to generate reports consisting of all or part of an individual patient's medical record (e.g. patient summary).	S.2.2	P			Report format may be plain text.
218				3. The system shall provide the ability to generate reports regarding multiple patients (e.g. diabetes roster).	S.2.2	N			Any disease registry might be included.
219				4. The system shall provide the ability to specify report parameters (sort and filter criteria) based on patient demographic and clinical data (e.g., all male patients over 50 that are diabetic and have a HbA1c value of over 7.0 or that are on a certain medication).	S.2.2	N			Minimum demographic data are age and gender.
220				5. The system shall provide the ability to access reports outside the EHR application.	S.2.2	P			For example, printed output, export to a file, etc.
221				6. The system shall provide the ability to produce reports based on the absence of a clinical data element (e.g., a lab test has not been performed or a blood pressure has not been measured in the last year).	S.2.2	N	N		
222				7. The system shall provide the ability to save report parameters for generating subsequent reports.	S.2.2	N			
224	F		Health record output: Allow users to define the records and/or reports that are considered the formal health record for disclosure purposes, and provide a mechanism for both chronological and specified record element output.	1. The system shall provide the ability to define one or more reports as the formal health record for disclosure purposes.	S.2.2.1	N			This allows the practice to not print demographics, certain confidential sections, or other items. Report format may be plain text initially. In the future there will be a need for structured reports as interoperability standards evolve.
225				2. The system shall provide the ability to generate hardcopy or electronic output of part or all of the individual patient's medical record.	S.2.2.1	P			This could include but is not limited to the ability to generate standardized reports needed for work, school, or athletic participation.


 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology					Revisions from prev. release (18SEP06) are in red text			Discussion / Comments
					Provisional Criteria (2007) are highlighted in yellow			
Original line # Phase I	WG	Category and Description	Specific Criteria	Source or References	Compliance Key:			
					Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
226			3. The system shall provide the ability to generate hardcopy and electronic output by activities and events on a chosen date and/or date range (e.g., all hospital discharge summaries) .	S.2.2.1	M			





AMBULATORY FUNCTIONALITY
Proposed 2007 Final Criteria - Feb 14 2007
Proposed FUNCTIONALITY For 2007 Certification of
Ambulatory EHRs incorporates AFWG work to 28 Nov 2006
 © 2007 The Certification Commission for Healthcare Information Technology

Revisions from prev. release (18SEP06) are in red text
 Provisional Criteria (2007) are highlighted in yellow
 Compliance Key:
 P = Previous Criteria
 N = New for Year
 M = Modified for Year

Original line # Phase I	WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
					Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
227			4. The system shall provide the ability to export structured data which removes those identifiers listed in the HIPAA definition of a limited dataset. This export on hardcopy and electronic output leaves the actual PHI data unmodified in the original record.	S.2.2.1	M		N	De-identifying data on hardcopy or electronic output is necessary for research. However, it must be emphasized that this function is not intended to cleanse the text in the note or data in the original record. As per HIPAA Standards for Privacy of Individually Identifiable Health Information, 45 CFR Parts 160 and 164, identifiers that shall be removed are: 1. Names; 2. Postal address information, other than town or city, state and zip code; 3. Telephone numbers; 4. Fax numbers; 5. Electronic mail addresses; 6. Social security numbers; 7. Medical record numbers; 8. Health plan beneficiary numbers; 9. Account numbers; 10. Certificate/license numbers; 11. Vehicle identifiers and serial numbers, including license plate numbers; 12. Device identifiers and serial numbers; 13. Web Universal Resource Locators (URLs); 14. Internet Protocol (IP) address numbers; 15. Biometric identifiers, including finger and voice prints; and 16. Full face photographic images and any comparable images.

Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
228				5. The system shall provide the ability to create hardcopy and electronic report summary information (procedures, medications, labs, immunizations, allergies, and vital signs).	S.2.2.1	P			The report that's produced should be organized by section to make it easier to read.
229				6. The system shall have the ability to provide support for disclosure management in compliance with HIPAA and applicable law.		N			This criterion may be satisfied by providing the ability to create a note in the patient's record. More advanced functionality may be market differentiators or requirements in later years.
230	F	Encounter management: Manage and document the health care delivered during an encounter.	1. The system shall provide the ability to document a patient encounter.	S.3.1	P				
231			2. The system shall provide the ability to document encounters by one or more of the following means: direct keyboard entry of text; structured data entry utilizing templates, forms, pick lists or macro substitution; dictation with subsequent transcription of voice to text, either manually or via voice recognition system.	S.3.1	P				This does not preclude entry via new technologies.
232			3. The system shall provide the ability to associate individual encounters with diagnoses.	S.3.1	P				
233			4. The system shall have the ability to provide filtered displays of encounters based on encounter characteristics, including date of service, encounter provider and associated diagnosis.	S.3.1	N				
234	F		Rules-driven financial and administrative coding assistance: Provide financial and administrative coding assistance based on the structured data available in the encounter documentation.	1. The system shall have the ability to provide a list of financial and administrative codes.	S.3.2.2	P			
235		2. The system shall provide the ability to select an appropriate CPT Evaluation and Management code based on data found in a clinical encounter.		S.3.2.2	P				May be accomplished via a link to another application.

Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
238	F	Eligibility verification and determination of coverage	1. The system shall provide the ability to display medical eligibility obtained from patient's insurance carrier, populated either through data entry in the system itself or through an external application interoperating with the system.	S.3.3.2	M			The EHR need only provide information for the physician as to whether the patient is covered by that insurance plan. This can be accomplished by a text note following telephone verification.	
240	F	Manage Practitioner/Patient relationships: Identify relationships among providers treating a single patient, and provide the ability to manage patient lists assigned to a particular provider.	1. The system shall provide the ability to identify by name all providers associated with a specific patient encounter.	S.3.4	P			A provider is defined as anyone delivering clinical care such as physicians, PAs, CNPs and nurses; the provider is the person who completes the note.	
242			3. The system shall provide the ability to specify the primary or principal provider responsible for the care of a patient within a care setting.		N				
244	F	Clinical decision support system guidelines updates: Receive and validate formatted inbound communications to facilitate updating of clinical decision support system guidelines and associated reference material	1. The system shall provide the ability to update the clinical content or rules utilized to generate clinical decision support reminders and alerts.	S.3.7.1	P			Growth charts, CPT-4 codes, drug interactions would be an example. Any method of updating would be acceptable. Content could be third part or customer created.	
245			2. The system shall provide the ability to update clinical decision support guidelines and associated reference material.	S.3.7.1	P			Any method of updating would be acceptable. Content could be third part or customer created.	
247		Enforcement of confidentiality: Enforce the applicable jurisdiction's patient privacy rules as they apply to various parts of an EHR-S through the implementation of security mechanisms.	1. The system shall provide the ability to audit the date/time and user of each instance when a patient chart is printed by the system .	I.1.9	N			Does not include screen print and other functions that are outside the EHR system.	
249			3. The system shall provide the ability to identify all users who have accessed an individual's chart over a given time period, including date and time of access .	I.1.9	N			Specific items/sections of information accessed shall be identified, with appropriate audit trail.	

Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
 <p>AMBULATORY FUNCTIONALITY Proposed 2007 Final Criteria - Feb 14 2007 Proposed FUNCTIONALITY For 2007 Certification of Ambulatory EHRs incorporates AFWG work to 28 Nov 2006 © 2007 The Certification Commission for Healthcare Information Technology</p>						Revisions from prev. release (18SEP06) are in red text Provisional Criteria (2007) are highlighted in yellow Compliance Key: P = Previous Criteria N = New for Year M = Modified for Year			
251				5. The system shall provide the ability to prevent specified user(s) from accessing a designated patient's chart	I.1.9	N	N		An example would be preventing access to a VIP or staff member's chart. When access is restricted, the system shall provide a means for appropriately authorized users to "break the glass" for emergency situations. Such overrides should be audited.
252	F	Data retention, availability, and destruction: Retain, ensure availability, and destroy health record information according to organizational standards. This	1. The system shall provide the ability to retain data until otherwise purged, deleted, archived or otherwise deliberately removed.	I.2.1	P				
257	F	Extraction of health record information: Manage data extraction in accordance with analysis and reporting requirements. The extracted data may require use of more than one application and it may be pre-processed (for example, by being de-identified) before transmission. Data extractions may be used to exchange data and provide reports for primary and ancillary purposes.	1. The system shall provide the ability to export (extract) pre-defined set(s) of data out of the system	I.2.4	N			For example, export of performance measures, ability to query data base, chronic disease management tools.	
258			2. The system shall provide the ability to import data into the system	I.2.4	N			Data import implies receiving discrete data into the EHR in an automated manner as opposed to manual data entry or document scanning. This could be accomplished via a real time or batch interface or a manual data load.	
259			3. The system shall provide the ability remove discrete patient identifiers.	I.2.4	N			De-identification is necessary for research purposes, e.g., to identify patterns of disease. External applications can be used to meet this criteria.	
261	F	Concurrent Use: EHR system supports multiple concurrent physicians through application, OS and database.	1. The system shall provide the ability for multiple users to interact concurrently with the EHR application.	Ontario 5.6.1.a	P				
262			2. The system shall provide the ability for concurrent users to simultaneously view the same record.	Ontario 5.6.1.a	P				

Original line # Phase I		WG	Category and Description	Specific Criteria	Source or References	Compliance			Discussion / Comments
						Certify in May 2007	Roadmap for May 2008	Roadmap for May 2009 and beyond	
263				3. The system shall provide the ability for concurrent users to view the same clinical documentation or template.	Ontario 5.6.1.a	P			
264				4. The system shall provide record-level protection to maintain the integrity of clinical data during concurrent access .	Ontario 5.6.1.a, I.1.9	P			To prevent users from simultaneously attempting to update a record with resultant loss of data