

Object Management Group

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Clinical Observations Access Service (COAS)

Request For Proposal (RFP)

OMG Document: corbamed/97-12-28

Submissions due: May, 18, 1998

Objective of this RFP

This RFP solicits proposals for accessing clinical observations. Clinical observations constitute a significant proportion of the information recorded about any patient. Examples of clinical observations include the following: laboratory results; vital signs; subjective and objective observations and assessments; observations and measurements provided by a specialist such as radiologist or pathologist who interprets images and other multi-media data. Interoperable specifications that support the activities involved in accessing clinical observations are sought in this RFP. The specifications should leverage existing standards such as HL7 and DICOM .

For further details see Chapter 6 of this document.

1.0 Introduction

1.1 Goals of OMG

The Object Management Group (OMG) is the world's largest software consortium with a membership of over 800 vendors, developers, and end users. Established in 1989, its mission is to promote the theory and practice of Object Technology (OT) for the development of distributed computing systems.

A key goal of OMG is create a standardized object-oriented architectural framework for distributed applications based on specifications that enable and support distributed objects. Objectives include the *reusability*, *portability*, and *interoperability* of object-oriented software components in heterogeneous environments. To this end, the OMG adopts interface and protocol specifications, based on commercially available object technology, that together define an Object Management Architecture (OMA).

1.2 Organization of this document

The remainder of this document is organized as follows:

Chapter 2 - *Architectural Context* - background information on OMG's Object Management Architecture.

Chapter 3 - *Adoption Process* - background information on the OMG specification adoption process.

Chapter 4 - *Instructions for Submitters* - explanation of how to make a submission to this RFP.

Chapter 5 - *General Requirements on Proposals* - requirements and evaluation criteria that apply to all proposals submitted to OMG.

Chapter 6 - *Specific Requirements on Proposals* - problem statement, scope of proposals sought, mandatory and optional requirements, issues to be discussed, evaluation criteria, and timetable that apply specifically to this RFP.

Additional RFP-specific chapters may also be included following Chapter 6.

1.3 References

The following documents are referenced in this document:

Richard Soley (ed.), *Object Management Architecture Guide*, Third Edition, Wiley, June 1995. OMG Document ab/97-05-05.

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The Common Object Request Broker: Architecture and Specification,
Revision 2.0, July 1996. OMG Document ptc/96-08-04.

CORBA services: Common Object Services Specification, Revised Edition,
March 1995. OMG Document formal/97-05-03.

CORBA facilities Architecture, Revision 4.0, November 1995.

Business Committee RFP Attachment, OMG Document omg/96-01-01.

Policies and Procedures of the OMG Technical Process, OMG Document
pp/97-06-01 or successor.

These documents can be obtained by contacting OMG at document@omg.org. Many OMG documents, including this document, are available electronically from OMG's document server. Send a message containing the single line "help" to server@omg.org for more information.

For more information about OMG visit OMG's Web page (URL <http://www.omg.org/>). If you have general questions about this RFP send email to responses@omg.org.

2.0 Architectural Context

2.1 Object Management Architecture

The *Object Management Architecture Guide* (OMAG) describes OMG's technical objectives and terminology and provides the conceptual infrastructure upon which supporting specifications are based. The guide includes the *OMG Object Model*, which defines common semantics for specifying the externally visible characteristics of objects in a standard implementation-independent way, and the *OMA Reference Model*.

The Reference Model identifies and characterizes the components, interfaces, and protocols that compose the OMA. This includes the Object Request Broker (ORB) component that enables clients and objects to communicate in a distributed environment, and four categories of object interfaces:

- *Object Services* are interfaces for general services that are likely to be used in any program based on distributed objects.
- *Common Facilities* are interfaces for horizontal end-user-oriented facilities applicable to most application domains.
- *Domain Interfaces* are application domain-specific interfaces.
- *Application Interfaces* are non-standardized application-specific interfaces.

A second part of the Reference Model introduces the notion of domain-specific *Object Frameworks*. An Object Framework component is a collection of cooperating objects that provide an integrated solution within an application or technology domain and which is intended for customization by the developer or user.

Through a series of RFPs, OMG is populating the OMA with detailed specifications for each component and interface category in the Reference Model. Adopted specifications include the Common Object Request Broker Architecture (CORBA), CORBAservices, and CORBAfacilities.

The wide-scale industry adoption of OMG's OMA provides application developers and users with the means to build interoperable software systems distributed across all major hardware, operating system, and programming language environments.

2.2 CORBA

The *Common Object Request Broker Architecture* defines the programming interfaces to the OMA ORB component. An ORB is the basic mechanism by which objects transparently make requests to - and receive responses from - each other on the same machine or across a network. A client need not be aware of the mechanisms used to communicate with or activate an object, how the object is implemented, nor where the object is located. The ORB thus forms the foundation for building applications constructed from distributed objects and for interoperability between applications in both homogeneous and heterogeneous environments.

The *OMG Interface Definition Language (IDL)* provides a standardized way to define the interfaces to CORBA objects. The IDL definition is the contract between the implementor of an object and the client. IDL is a strongly typed declarative language that is programming language-independent. Language mappings enable objects to be implemented and sent requests in the developer's programming language of choice in a style that is natural to that language.

CORBA 2.0 is an extension and restructuring of the earlier CORBA 1.2 specification. CORBA 2.0 is a family of specifications consisting of the following components:

- Core (including IDL syntax and semantics)
- Interoperability
- An expanding set of language mappings, including:

C

C++

SmallTalk

Ada95

COBOL

Each component is a separate compliance point. The minimum required for a CORBA-compliant implementation is adherence to the core and one language mapping.

2.3 CORBA/Interoperability

Interoperability between CORBA-compliant ORBs is provided by OMG's *Internet Inter-ORB Protocol (IIOP)*. Adopted in December 1994 as the

mandatory CORBA 2.0 protocol for “out of the box” interoperability, IIOP is the TCP/IP transport mapping of a *General Inter-ORB Protocol (GIOP)*. IIOP enables requests to be sent to networked objects managed by other ORBs in other domains.

The OMG interoperability architecture also accommodates communication using optional *Environment-Specific IOPs (ESIOPs)*, the first of which is the DCE-CIOP.

2.4 CORBA services

Object Services are general purpose services that are either fundamental for developing useful CORBA-based applications composed of distributed objects, or that provide a universal - application domain-independent - basis for application interoperability.

Object Services are the basic building blocks for distributed object applications. Compliant objects can be combined in many different ways and put to many different uses in applications. They can be used to construct higher level facilities and object frameworks that can interoperate across multiple platform environments.

Adopted OMG Object Services are collectively called CORBA services and include Naming, Events, LifeCycle, Persistent Object, Relationships, Externalization, Transactions, Concurrency Control, Licensing, Query, Properties, Security, Time, Collections, and Trading Services.

2.5 CORBA facilities

Common Facilities are interfaces for horizontal end-user-oriented facilities applicable to most domains. Adopted OMG Common Facilities are collectively called CORBA facilities and include an OpenDoc-based Distributed Document Component Facility.

A specification of a Common Facility or Object Service typically includes the set of interface definitions - expressed in OMG IDL - that objects in various roles must support in order to *provide, use, or participate in* the facility or service. As with all specifications adopted by OMG, facilities and services are defined in terms of interfaces and their semantics, and not a particular implementation.

2.6 Object Frameworks and Domain Interfaces

Unlike the interfaces to individual parts of the OMA “plumbing” infrastructure, Object Frameworks are complete higher level components that provide

functionality of direct interest to end-users in particular application or technology domains. They are vertical slices down the OMG “interface stack”.

Object Frameworks are collections of cooperating objects categorized into *Application, Domain, Facility, and Service Objects*. Each object in a framework supports (through interface inheritance) or makes use of (via client requests) some combination of Application, Domain, CORBAfacilities, and CORBAservices *interfaces*.

A specification of an Object Framework defines such things as the structure, interfaces, types, operation sequencing, and qualities of service of the objects that make up the framework. This includes requirements on implementations in order to guarantee application portability and interoperability across different platforms.

Domain Task Force RFPs are likely to focus on Object Framework specifications that include new Domain Interfaces for application domains such as Finance, Healthcare, Manufacturing, Telecom, Electronic Commerce, and Transportation.

3.0 Adoption Process

3.1 Introduction

OMG adopts specifications for interfaces and protocols by explicit vote on a technology-by-technology basis. The specifications selected each fill in a portion of the OMA Reference Model. OMG bases its decisions on both business and technical considerations. Once a specification is adopted by OMG, it is made available for use by both OMG members and non-members.

For more detailed information on the adoption process see the *Policies and Procedures of the OMG Technical Process*.

3.2 Role of Board of Directors

The OMG Board of Directors votes to formally adopt specifications on behalf of OMG. The OMG Technology Committees (Domain and Platform TCs) and Architecture Board (AB) provide technical guidance to the Board of Directors. In addition, the Business Committee of the Board provides guidance to ensure that implementations of adopted specifications are made commercially available.

3.3 Role of Technology Committees and Architecture Board

Submissions to RFPs are evaluated by the TC Task Force (TF) that initiated the RFP. Selected specifications are recommended to the parent TC after being reviewed by the Architecture Board for consistency with the OMA. The full TC then votes to *recommend adoption* to the OMG Board.

3.4 Role of Task Forces

The role of the initiating TF is to technically evaluate submissions and select one or more specifications that satisfy the requirements of the RFP. The process typically takes the following form:

- Voter Registration

Interested TF members may register to participate in specification selection votes for an RFP. Registration ends on a specified date 6 or more weeks after the announcement of the registration period. The registration closure date is typically around the time of initial submissions. Companies who have submitted an LOI are automatically registered to vote.

- Initial Submissions

Initial submissions are due by a specified deadline. Submitters normally present their proposals at the next following meeting of the TF. Initial submissions are expected to be full and complete proposals and working implementations of the proposed specifications are expected to exist at the time of submission.

- Evaluation Phase

A period of approximately 120 days follows during which the TF evaluates the submissions. During this time submitting companies have the opportunity to revise and/or merge their initial submissions, if they so choose.

- Revised Submissions

Final revised submissions are due by a specified deadline. Submitters again normally present their proposals at the next following meeting of the TF. Finalists may be requested to demonstrate implementations of their proposal.

- Selection Vote

When the registered voters of the TF believe that they sufficiently understand the relative merits of the revised submissions, a specification selection vote is taken.

3.5 Goals of the evaluation

The primary goals of the TF evaluation process are to:

- Provide a fair and open process
- Force a critical review of the submissions and discussion by all members of the TF
- Give feedback to allow submitters to address concerns in their revised submissions
- Build consensus on acceptable solutions
- Enable voting members to make an informed selection decision

Submitters are expected actively to contribute to the evaluation process.

4.0 Instructions for Submitters

4.1 OMG Membership

Submissions to this RFP may only be made by Contributing or Domain Contributing members of the OMG; to submit to an RFP issued by the Platform Technology Committee an organisation must be a Contributing member at the date of the submission deadline, while for Domain Technology RFPs the submitter or submitters must be either Contributing or Domain Contributing members. Submitters sometimes choose to name other organisations that support a submission in some way; however, this has no formal status within the OMG process, and for OMG's purposes confers neither duties nor privileges on the organisations concerned.

4.2 Submission Effort

Unlike a submission to an OMG Request For Information (RFI), an RFP submission may require significant effort in terms of document preparation, presentations to the initiating TF, and participation in the TF evaluation process. Several staff months of effort might be necessary. OMG is unable to reimburse submitters for any costs in conjunction with their submissions to this RFP.

4.3 Letter of Intent

A Letter of Intent (LOI) must be submitted to the OMG Business Committee signed by an officer of your organization signifying your intent to respond to the RFP and confirming your organization's willingness to comply with OMG's terms and conditions, and commercial availability requirements. These terms, conditions, and requirements are defined in the *Business Committee RFP Attachment* and are reproduced verbatim in section 4.3 below.

The LOI should designate a single contact point within your organization for receipt of all subsequent information regarding this RFP and your submission. The name of this contact will be made available to all OMG members. The LOI is typically due 60 days before the deadline for initial submissions. LOIs must be sent by fax or paper mail to the "RFP Submissions Desk" at the main OMG address shown on the first page of this RFP.

Here is a suggested template for the Letter of Intent:

This letter confirms the intent of <__organisation required__> (the organisation) to submit a response to the OMG <__RFP name required__> RFP. We will grant OMG and its members the right to copy our response for

review purposes as specified in section 4.6 of the RFP. Should our response be adopted by OMG we will comply with the OMG Business Committee terms set out in section 4.3 of the RFP and in document omg/97-01-01.

<____contact name and details required____> will be responsible for liaison with OMG regarding this RFP response.

The signatory below is an officer of the organisation and has the approval and authority to make this commitment on behalf of the organisation.

<____signature required____>

4.4 Business Committee RFP Attachment

This section contains the text of the Business Committee RFP attachment concerning commercial availability requirements placed on submissions. This attachment, available separately as document omg/97-10-01, was approved by the OMG Board in September 1997.

Commercial considerations in OMG technology adoption

A1 Introduction

OMG wishes to encourage rapid commercial adoption of the specifications it publishes. To this end, there must be neither technical, legal nor commercial obstacles to their implementation. Freedom from the first is largely judged through technical review by the relevant OMG Technology Committee; the second two are the responsibility of the OMG Business Committee. The BC also looks for evidence of a commitment by a submitter to the commercial success of products based on the submission.

A2 Business Committee evaluation criteria

A2.1 Viable to implement across platforms

While it is understood that final candidate OMG submissions often combine technologies before they have all been implemented in one system, the Business Committee nevertheless wishes to see evidence that each major feature has been implemented, preferably more than once, and by separate organisations. Pre-product implementations are acceptable. Since use of OMG specifications should not be dependant on any one platform, cross-platform availability and interoperability of implementations should be also be demonstrated.

A2.2 Commercial availability

In addition to demonstrating the existence of implementations of the specification, the submitter must also show that products based on the specification are commercially available, or will be within 12 months of the date when the specification was recommended for adoption by the appropriate Task Force. Proof of intent to ship product within 12 months might include:

- *A public product announcement with a shipping date within the time limit.*
- *Demonstration of a prototype implementation and accompanying draft user documentation.*

Alternatively, and at the Business Committee's discretion, submissions may be adopted where the submitter is not a commercial software provider, and therefore will not make implementations commercially available. However, in this case the BC will require concrete evidence of two or more independent implementations of the specification being used by end-user organisations as part of their businesses.

Regardless of which requirement is in use, the submitter must inform the OMG of completion of the implementations when commercially available.

A2.3 Unencumbered technology

The submitter must certify that it is not aware of any claim that the specification infringes any patent, copyright, or other intellectual property right (collectively referred to in this policy statement as "IPR"). Except for this certification, the submitter will not be required to make any other warranty, and specifications will be offered by OMG for implementation "as is". If the submitter owns IPR to which an implementation of a specification based upon its submission would necessarily be subject, it must certify to the BC that it will make a suitable license available to any implementer to permit development and commercialisation of an implementation without violation of such IPR.

The submitter shall also certify that this license will be made available on commercially reasonable, non-discriminatory terms. The submitter is responsible for disclosing in detail all known restrictions and fees, placed either by the submitter or others, on technology necessary for implementation of the specification.

A2.4 Publication of the specification

Should the submission be adopted, the submitter must grant OMG (and its sublicensees) a world-wide, royalty-free licence to edit, store, duplicate and distribute both the specification and works derived from it (such as revisions

and teaching materials). This requirement applies only to the written specification, not to any implementation of it.

A2.5 Continuing support

The submitter must show a commitment to continue supporting the technology underlying the specification after OMG adoption, for instance by showing the BC development plans for future revisions, enhancement or maintenance.

4.5 Responding to RFP items

4.5.1 Separate proposals

Unless otherwise indicated in Chapter 6, independent proposals are solicited for each separate item in the RFP. Each item is considered a separate architectural entity for which a proposal may be made. A submitter may respond to any or all items. Each item will be evaluated independently by the initiating TF.

Submissions that do not present clearly separable proposals for multiple items may therefore be at a disadvantage.

It should be noted that a given technology (e.g. software product) may support two or more RFP items. So long as the interfaces for each item are separable, this is not precluded.

4.5.2 Complete proposals

Proposals for each separate RFP item must be complete. A submission must propose full specifications for each item and address all the relevant general and specific requirements detailed in this RFP.

4.5.3 Additional specifications

Submissions may include additional specifications for items not covered by the RFP which they believe to be necessary and integral to their proposal. Information on these additional items should be clearly distinguished.

Submitters must give a detailed rationale as to why these specifications should also be considered for adoption. However submitters should note that a TF is unlikely to consider additional items that are already on the roadmap of an OMG TF, since this would preempt the normal adoption process.

4.5.4 Alternative approaches

Submitters may provide alternative RFP item definitions, categorizations, and groupings so long as the rationale for doing so is clearly stated. Equally, submitters may provide alternative models for how items are provided within the OMA if there are compelling technological reasons for a different approach.

4.6 Confidential and Proprietary Information

The OMG specification adoption process is an open process. Responses to this RFP become public documents of the OMG and are available to members and non-members alike for perusal. No confidentiality or proprietary information of any kind will be accepted in a submission to this RFP.

4.7 Copyright Waiver

If a submitted document is copyrighted, a waiver of copyright for unlimited duplication by the OMG is required to be stated in the document. In addition, a limited waiver of copyright is required that allows each OMG member to make up to fifty (50) copies of the document for review purposes only.

4.8 Proof of Concept

Submissions must include a “proof of concept” statement, explaining how the submitted specifications have been demonstrated to be technically viable. The technical viability has to do with the state of development and maturity of the technology on which a submission is based. This is not the same as commercial availability. Proof of concept statements can contain any information deemed relevant by the submitter, for example:

“This specification has completed the design phase and is the process of being prototyped.”

“An implementation of this specification has been in beta-test for 4 months.”

“A named product (with a specified customer base) is a realization of this specification.”

It is incumbent upon submitters to demonstrate to the satisfaction of the TF the technical viability of their proposal. OMG will favor proposals based on technology for which sufficient relevant experience has been gained in CORBA-based or comparable environments.

4.9 Format of RFP Submissions

This section provides guidance on how to structure your RFP submission.

4.9.1 General

- Submissions that are concise and easy to read will inevitably receive more consideration.
- Submitted documentation should be confined to that directly relevant to the items requested in the RFP. If this is not practical, submitters must make clear what portion of the documentation pertains directly to the RFP and what portion does not.
- The models and terminology in the *Object Management Architecture Guide* and *CORBA* should be used in your submission. Where you believe this is not appropriate, describe and provide a rationale for the models and terminology you believe OMG should use.

4.9.2 Suggested Outline

A three part structure for submissions is suggested:

PART I

- Copyright Waiver (see 4.5)
- Submission contact point (see 4.2)
- Overview or guide to the material in the submission
- Overall design rationale (if appropriate)
- Statement of proof of concept (see 4.6)
- Resolution of RFP mandatory and optional requirements

Explain how your proposal satisfies the mandatory and (if applicable) optional requirements stated in Chapter 6. References to supporting material in Part II should be given.

In addition, if your proposal does not satisfy any of the general requirements stated in Chapter 5, provide a detailed rationale.

- Responses to RFP issues to be discussed

Discuss each of the “Issues To Be Discussed” identified in Chapter 6.

PART II

- Proposed specification

PART III

- Summary of optional versus mandatory interfaces

Submissions must clearly distinguish interfaces that all implementations must support from those that may be optionally supported.

- Proposed compliance points

Submissions should propose appropriate compliance points for implementations.

- Changes or extensions required to adopted OMG specifications

Submissions must include a full specification of any changes or extensions required to existing OMG specifications. This should be in a form that enables “mechanical” section-by-section revision of the existing specification.

- Complete IDL definitions

For reference purposes and to facilitate electronic usage, submissions should reproduce in one place a complete listing in compilable form of the IDL definitions proposed for standardization.

4.10 How to Submit

Submitters should send an electronic version of their submission to the *RFP Submissions Desk* (rfp@omg.org) at OMG by 5:00 PM U.S. Eastern Standard Time (22:00 GMT) on the day of the submission deadline. Acceptable formats are Postscript, ASCII, PDF, FrameMaker, Word, and WordPerfect. However, it should be noted that a successful submission must be supplied to OMG’s technical editors in Framemaker source format, using the most recent available OMG submission template (document ab/96-06-02 at the time of writing). The AB will not endorse adoption of any submission for which appropriately-formatted Framemaker sources are not available; it may therefore be convenient to prepare all stages of a submission using this template.

Submitters should make sure they receive electronic or voice confirmation of the successful receipt of their submission. Submitters should also send, within three (3) working days after the submission deadline, a single hardcopy version of their submission to the attention of the “RFP Submissions Desk” at the main OMG address shown on the first page of this RFP.

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In addition, submitters are responsible for making available 100 paper copies to attendees of the TF meeting immediately following a submission deadline. There are normally two such presentation meetings, one for the initial and one for the revised submissions.

5.0 General Requirements on Proposals

5.1 Mandatory Requirements

- 5.1.1 Proposals shall express interfaces in OMG IDL. Proposals should follow accepted OMG IDL and CORBA programming style. The correctness of the IDL shall be verified using at least one IDL compiler (and preferably more than one). In addition to IDL quoted in the text of the submission, all the IDL associated with the proposal shall be supplied to OMG in compiler-readable form.
- 5.1.2 Proposals shall specify *operation behavior*, *sequencing*, and *side-effects* (if any).
- 5.1.3 Proposals shall be *precise* and *functionally complete*. There should be no implied or hidden interfaces, operations, or functions required to enable an implementation of the proposed specification.
- 5.1.4 Proposals shall clearly distinguish *mandatory* interfaces and other specification elements that all implementations must support from those that may be *optionally* supported.
- 5.1.5 Proposals shall *reuse* existing OMG specifications including CORBA, CORBA services, and CORBA facilities in preference to defining new interfaces to perform similar functions.
- 5.1.6 Proposals shall justify and fully specify any *changes or extensions* required to existing OMG specifications. This includes changes and extensions to CORBA inter-ORB protocols necessary to support interoperability. In general, OMG favors *upwards compatible* proposals that minimize changes and extensions to existing OMG specifications.
- 5.1.7 Proposals shall factor out functions that could be used in different contexts and specify their interfaces separately. Such *minimality* fosters re-use and avoids functional duplication.
- 5.1.8 Proposals shall use or depend on other interface specifications only where it is actually necessary. While re-use of existing interfaces to avoid duplication will be encouraged, proposals should avoid gratuitous use.

- 5.1.9 Proposals shall specify interfaces that are *compatible* and can be used with existing OMG specifications. Separate functions doing separate jobs should be capable of being used together where it makes sense for them to do so.
- 5.1.10 Proposals shall preserve maximum *implementation flexibility*. Implementation descriptions should not be included, however proposals may specify constraints on object behavior that implementations need to take into account over and above those defined by the interface semantics.
- 5.1.11 Proposals shall allow *independent implementations* that are *substitutable* and *interoperable*. An implementation should be replaceable by an alternative implementation without requiring changes to any client.
- 5.1.12 Proposals shall be compatible with the architecture for system distribution defined in ISO/IEC 10746, Reference Model of Open Distributed Processing (ODP). Where such compatibility is not achieved, the response to the RFP must include reasons why compatibility is not appropriate and an outline of any plans to achieve such compatibility in the future.
- 5.1.13 In order to demonstrate that the service or facility proposed in response to this RFP, can be made secure in environments requiring security, answers to the following questions shall be provided:
- What, if any, are the security sensitive objects that are introduced by the proposal?
 - Which accesses to security-sensitive objects must be subject to security policy control?
 - Does the proposed service or facility need to be security aware?
 - What CORBAsecurity level and options are required to protect an implementation of the proposal? In answer to this question, a reasonably complete description of how the facilities provided by the level and options (e.g. authentication, audit, authorization, message protection etc.) are used to protect access to the sensitive objects introduced by the proposal shall be provided.
 - What default policies should be applied to the security sensitive objects introduced by the proposal?
 - Of what security considerations must the implementers of your proposal be aware?

5.2 Evaluation criteria

Although the OMG adopts interface specifications, the technical viability of implementations will be taken into account during the evaluation process. The following criteria will be used:

5.2.1 Performance

Potential implementation trade-offs for performance will be considered.

5.2.2 Portability

The ease of implementation on a variety of ORB systems and software platforms will be considered.

5.2.3 Compliance: Inspectability and Testability

The adequacy of proposed specifications for the purposes of compliance inspection and testing will be considered. Specifications should provide sufficient constraints on interfaces and implementation characteristics to ensure that compliance can be unambiguously assessed through both manual inspection and automated testing.

6.0 Specific Requirements on Proposals

This RFP solicits proposals for accessing clinical observations. Clinical observations constitute a significant proportion of the information recorded about any patient. Examples of clinical observations include the following: laboratory results; vital signs; subjective and objective observations and assessments; observations and measurements provided by a specialist such as radiologist or pathologist who interprets images and other multi-media data. Interoperable specifications that support the activities involved in accessing clinical observations are sought in this RFP. The specifications should leverage existing standards such as HL7 and DICOM.

6.1 Problem Statement

Throughout this RFP the concept “Clinical Observations” indicates any information that has been captured about a single patient’s medical/physical state and relevant context information. This may be derived by instruments such as in the case of images, vital signs and lab results or it may be derived by a health professional via direct examination of the patient and transcribed. This term applies to information that has been captured whether or not it has been reviewed by an appropriate authority to confirm its applicability to the patient record. The term is used independently of any particular information representation format.

Accessing information from current clinical observations systems is not easy. They exist in heterogeneous systems and require custom and often proprietary mechanisms to access this information. There is a lack of a coherent way to access information from the various systems that can provide clinical observations. Even when existing systems support standards, these systems do not interoperate for a number of reasons. The semantics are often times ambiguous, or there exists multiple overlapping standards that contain subsets of clinical observations. There is a clear need to provide a uniform and standard way to access clinical observations, the focus of this RFP.

6.2 Scope of Proposals Sought

Proposals are sought that provide clear, unambiguous, mechanisms to support accessing a wide variety of clinical observations.

6.3 Relationship to Existing OMG Specifications

Technology submissions should take advantage of existing OMG specifications where appropriate. The following services may be appropriate:

- Security
- Time
- Trader
- Event

In addition, submitters should be aware that RFP's are currently outstanding for technology adoption in the following areas. Submitters are encouraged to follow this work, because technology may be adopted in this space before submissions are due for this RFP. These services include:

- Notifications
- Messaging
- Objects by value
- Patient Identification Service (PIDS)
- Lexicon Query Service (LQS)
- Workflow
- Component Model
- Tagged service.

Future CORBAMED RFPs are anticipated that will use the COAS such as Decision Support.

6.4 Related Documents and Standards

Some groundwork in this area has already been laid by other standards organizations, particularly: DICOM, HL7, GEHR, IEEE 1073 MIB, ASTM and CEN TC 251.

The following are the responses to the Clinical Observations RFI issued by CORBAMED in June 1997. Submitters should reference these for important evaluation criteria that may be applied by members of the CORBAMED task force.

- corbamed/97-08-07, 97-08-06, 97-08-05 (CHIME and The GEHR Architecture)

- corbamed/97-09-05 (Joint Response from Philips Medical Systems, Baptist Health Systems or South Florida, CareFlow|Net, Inc. and Kurzweil)
- corbamed/97-08-08 (Yale University School of Medicine)
- corbamed/97-08-09, 97-08-04, 97-08-10 (Protocol Systems and University of Michigan Health Services)
- corbamed/97-09-04 (HL7 SGML/XML SIG)
- corbamed/97-09-06 (American Association For Medical Transcription)
- corbamed/97-09-07 (DICOM Working Group 8)
- corbamed/97-09-08 (HL7 IMSIG)

In addition the following white paper may be of use: corbamed/97-01-01 (white paper edited by Rachael Sokolowski). All of these documents are available from the OMG CORBAmed web site: <http://www.omg.org/docs/corbamed>.

6.5 Mandatory Requirements

This RFP solicits proposals for OMG IDL specifications of a Healthcare Clinical Observations Access Service (COAS) that will facilitate the access of clinical observations.

The following are the mandatory requirements for COAS:

- The COAS submission shall allow clinical observations to be requested, and/or sent.
- The COAS submission shall provide for filtering, such as by patient, provider, observation type, state and/or time frame. In addition to returning the observations, the COAS submission shall provide a mechanism for querying which instances of observations are available.
- The COAS submission shall support access to context information associated with observations.
- The COAS shall support a wide variety of observation types that can be extended by individual implementations and standardization. The COAS submission shall define a default set of clinical observations data types, for example: waveforms, text, images, audio, video, etc.

- The COAS submission shall not the preclude the use of publicly available vocabulary standards.

6.6 Optional Requirements

The following are optional requirements that a submission may decide to address but are not required to.

- The COAS submission may support a publish/subscribe mechanism where clients can to be notified when certain observations are available, have changed or meet a particular criteria. Subscriptions to events may be able to be controlled by entities separate from the receiver of the event. Submissions shall leverage the Notification or Event Service for this.
- The COAS submission may provide a mechanism for implementations to expose which types of observations and data they support in order to enable dynamic discovery of capabilities.
- The COAS submission may provide a general filtering mechanism for querying clinical observations.
- The COAS submission may define Trader service types to be used in registering with a Trader.
- COAS submission may support mechanisms to access historical changes made to observations.
- Submitters may provide an information model and an explicit mapping to their IDL. The information model shall be consistent with an industry accepted standard.
- The COAS submission may utilize closed, proprietary vocabularies *in addition* to the publicly-available vocabularies.
- The COAS submission may enable the use of OMG Lexicon Query Service (LQS) to support standard and non-standard vocabularies for Clinical Observations.

6.7 Issues to be discussed

If applicable, the COAS submission shall address the following:

- Describe possible approaches to integrate the use of the submission with HL7 and DICOM capabilities.

- Does the submission differentiate between clinical observations and other aspects of patient records? Describe why they should or should not be considered different.
- Proposal should discuss clinical context information management including: linkages to orders, encounters, and problems. For example submissions may discuss the linkage of pending orders and clinical observations.
- Describe the relationship between the Audio/Video Streams specification and the transmission of audio and video data.

6.8 Evaluation Criteria

The proposals will be evaluated on their completeness and their ability to address the mandatory requirements. A submitter should give justification for any mandatory requirements not met. In particular the following will be some of the criteria used in evaluating responses.

- Can the submission be used for a large variety of healthcare information systems that deal with clinical observations? Does this include Electronic Medical Record (EMR) systems?
- Does the submission facilitate the communications of clinical observations that is needed for workflow pertaining to capturing, managing, reporting, dispatching, signing, notification, and verification?
- Is the submission useful for both intra-enterprise and inter-enterprise communications of clinical observations?
- Is the submission independent of whether the integration of patient data is done at desktop applications, by a back end server, or a Clinical Data Repository (CDR)? Does it preclude any of these?
- Is the submission useful for a wide variety of business processes that handle clinical observations?
- Does the event subscription part of the submission inter-operate with Event Services specification, and/or the Notification Service?
- Does the submission reuse other OMG specifications where appropriate? In particular does it use or at least is it orthogonal to the Person Identification Service (PIDS) and Lexicon Query Service (LQS)?

- Can the submission be used to create services that can be efficient where bandwidth limitations exist? Can it be used to for real time applications? Can it be used for continuous data channels?
- Degree to which submission leverages existing standards.

6.9 RFP Timetable

The timetable for this RFP is given below. Note that the TF may, in certain circumstances, extend deadlines while the RFP is running, or may elect to have more than one revised submission step. The latest timetable can always be found in the Member Services section of OMG's Web page (URL <http://www.omg.org/>).

Event or Activity	Actual Date
TC votes to issue RFP	December 5, 1997
LOI to submit to RFP due	April 3, 1998
Voter registration closes	June 1, 1998
Initial submissions due	May 18, 1998
Revised submissions due	August 24, 1998
TF votes to select specifications	September 17, 1998
TC votes to recommend specifications	September-October, 1998
BOD votes to adopt specifications	November, 1998