

IQPP Corporate Audit Report Form and Checklist

Version 7.0 Effective April 1, 2015





IQPP Corporate Audit Report Form Version 7.0

Government Authority Identification	 [] For Certification/Recertification, pending resolution of issues listed on report form Section(s)Page(s) [] Significant issues listed on report form, Section(s)/Page(s) [] Recommend Re-audit within days. 	
Telephone Fax Person Responsible for Quality Assurance	 [] For Certification/Recertification, pending resolution of issues listed on report form Section(s)Page(s) [] Significant issues listed on report form, Section(s)/Page(s) 	
Telephone Fax Person Responsible for Quality Assurance	 [] For Certification/Recertification, pending resolution of issues listed on report form Section(s)Page(s) [] Significant issues listed on report form, 	
Telephone	[] For Certification/Recertification, pending resolution of issues listed on report form Section(s)Page(s)	,
Telephone	[] For Certification/Recertification, pending resolution of issues listed on report form	,
Telephone Fax Person Responsible for Quality Assurance		,
Telephone Fax Person Responsible for Quality Assurance	Auditor Recommendation: [] For Certification/Recertification	n.
Telephone Fax Person Responsible for Quality Assurance		
TelephoneFax Person Responsible for Quality Assurance Date of auditStart Time (approx.) End Time Other regulatory and government-agency approvals		
Telephone Fax Person Responsible for Quality Assurance		
Telephone Fax Person Responsible for Quality Assurance Date of audit Date of audit Start Time	Other regulatory and government-agency approvals	
Telephone Fax Person Responsible for Quality Assurance	(approx.) End Time	
TelephoneFax	Date of auditStart Time	
	Person Responsible for Quality Assurance	
	TelephoneFax	
AuditorCompany	Address	





Auditor's Statement

As an Auditor for the International Quality Plasma Program (IQPP), I shall not, either directly or indirectly, for myself or for the benefit of or in conjunction with any other person, corporation, partnership, association, agency, department, or other legal entity, use, communicate or otherwise disclose, or permit to be disclosed, any Confidential Information relating to this audit or company without prior written consent of such company; provided, however, Auditor may, only to the extent reasonably necessary or appropriate to the performance of Auditor's duties, disclose such Confidential Information to PPTA or an employee of PPTA for use in the IQPP Certification or a person to whom disclosure is otherwise required by applicable state or federal law or regulation.

All information obtained during audit will be forwarded to PPTA to be made a part of the company's IQPP certification file.

As a consultant appointed by PPTA to perform this company's IQPP audit, I hereby attest that to the best of my knowledge no conflict of interest exists between my current clients and the audited company and/or PPTA.

As a consultant for the purposes of performing the IQPP audit of said company, I certify that the attached audit findings and comments are true and accurate findings based on my observations and record review during the audit.

Auditor Signature	Date

POST AUDIT REVIEW

I acknowledge that the Auditor has reviewed the observations listed in this report. My signature does not constitute concurrence or denial of any of the observations made by the Auditor.

Company Representative_____Date_____

Title_____Center Name/Location _____





IQPP Corporate Audit Checklist Version 7.0

Α-	Qualified Donors, Donor Record File (DRF) Review &	Donoi	[.] Priv	асу
#	Audit Question	Yes	No	Ranking
1.	Does the company have written procedures or, in the			Critical
	case of automated donor process, functional			
	documentation (specifications/validation) that conform			
	to the IQPP Qualified Donor Standard?			
2.	Does the company have written procedures to track			Critical
	Applicant Donor Units (orphan units) as to their final			
	disposition?			
<u>Auc</u>	ditor Comments on Section A:			
В-	Community-Based Donor Population			
	Community-Based Donor Population	Yes	No	Ranking
#	Audit Question	Yes	No	Ranking
	Audit Question Does the company have written procedures in place	Yes	No	Ranking Critical
#	Audit Question Does the company have written procedures in place that conform to the IQPP Community-Based Donor	Yes	No	
# 1.	Audit Question Does the company have written procedures in place that conform to the IQPP Community-Based Donor Standard?	Yes	No	
# 1.	Audit Question Does the company have written procedures in place that conform to the IQPP Community-Based Donor	Yes	No	
# 1.	Audit Question Does the company have written procedures in place that conform to the IQPP Community-Based Donor Standard?	Yes	No	
# 1.	Audit Question Does the company have written procedures in place that conform to the IQPP Community-Based Donor Standard?	Yes	No	
# 1.	Audit Question Does the company have written procedures in place that conform to the IQPP Community-Based Donor Standard?	Yes	No	
# 1.	Audit Question Does the company have written procedures in place that conform to the IQPP Community-Based Donor Standard?	Yes	No	
# 1.	Audit Question Does the company have written procedures in place that conform to the IQPP Community-Based Donor Standard?	Yes	No	
# 1.	Audit Question Does the company have written procedures in place that conform to the IQPP Community-Based Donor Standard?	Yes	No	
# 1.	Audit Question Does the company have written procedures in place that conform to the IQPP Community-Based Donor Standard?	Yes	No	
# 1.	Audit Question Does the company have written procedures in place that conform to the IQPP Community-Based Donor Standard?	Yes	No	
# 1.	Audit Question Does the company have written procedures in place that conform to the IQPP Community-Based Donor Standard?	Yes	No	
# 1.	Audit Question Does the company have written procedures in place that conform to the IQPP Community-Based Donor Standard?	Yes	No	



C – Use of the National Donor Deferral Registry or centralized donor deferral registry usage				
#	Audit Question	Yes	No	Ranking
1.	Does the company have written procedures in place that conform to the IQPP Use of the NDDR Standard?			Critical
	<u>ditor Comments on Section C</u> :			
D -	Donor Education			
#	Audit Question	Yes	No	Ranking
1.	Does the company have written procedures in place that conform to the IQPP Donor Education Standard?			Major
2.	Can the company show evidence of donor education material that is provided to the plasma centers to educate donors?			Minor
<u>Au</u>	ditor Comments on Section D:			





F_	Personnel Education and Training			
#	Audit Question	Yes	No	Ranking
1.	Does the company have a written training guide with			Major
	instructions or procedures to be performed by the			
	trainee for each relevant plasma center job function?			
2.	Does the company have an annual GMP training policy			Major
	for all plasma center employees?			
3.	Is there a policy and process in place to verify that			Minor
	plasma center employees (with a functional job related			
	to donor screening, plasma collection, product handling			
	or other similar functions) have attained the minimum			
	level of education required in the Standard?			
Auc	ditor Comments on Section E:			
F-	Professional Plasma Collection Facility			
		licable	for th	e Corporate
	TE: There are no questions from this section that are appl	licable	for th	e Corporate
NO	TE: There are no questions from this section that are appl	licable	for th	e Corporate
NO Auc	TE: There are no questions from this section that are appl dit.	licable	for th	e Corporate
NO Auc G -	TE: There are no questions from this section that are app dit. • Complaints			
NO Auc G - #	TE: There are no questions from this section that are appl dit. - Complaints Audit Question	licable Yes	for th	Ranking
NO Auc G -	TE: There are no questions from this section that are appl dit. - Complaints Audit Question Does the company have written procedures available			
NO Auc G - #	TE: There are no questions from this section that are app dit. - Complaints Audit Question Does the company have written procedures available for receiving, recording and evaluating customer and/or			Ranking
NO Auc G - #	TE: There are no questions from this section that are appl dit. - Complaints Audit Question Does the company have written procedures available			Ranking
NO Auc G - # 1.	TE: There are no questions from this section that are appldit. - Complaints Audit Question Does the company have written procedures available for receiving, recording and evaluating customer and/or donor complaints?			Ranking
NO Auc G - # 1.	TE: There are no questions from this section that are app dit. - Complaints Audit Question Does the company have written procedures available for receiving, recording and evaluating customer and/or			Ranking
NO Auc G - # 1.	TE: There are no questions from this section that are appldit. - Complaints Audit Question Does the company have written procedures available for receiving, recording and evaluating customer and/or donor complaints?			Ranking
NO Auc G - # 1.	TE: There are no questions from this section that are appldit. - Complaints Audit Question Does the company have written procedures available for receiving, recording and evaluating customer and/or donor complaints?			Ranking
NO Auc G - # 1.	TE: There are no questions from this section that are appldit. - Complaints Audit Question Does the company have written procedures available for receiving, recording and evaluating customer and/or donor complaints?			Ranking
NO Auc G - # 1.	TE: There are no questions from this section that are appldit. - Complaints Audit Question Does the company have written procedures available for receiving, recording and evaluating customer and/or donor complaints?			Ranking
NO Auc G - # 1.	TE: There are no questions from this section that are appldit. - Complaints Audit Question Does the company have written procedures available for receiving, recording and evaluating customer and/or donor complaints?			Ranking
NO Auc G - # 1.	TE: There are no questions from this section that are appldit. - Complaints Audit Question Does the company have written procedures available for receiving, recording and evaluating customer and/or donor complaints?			Ranking
NO Auc G - # 1.	TE: There are no questions from this section that are appldit. - Complaints Audit Question Does the company have written procedures available for receiving, recording and evaluating customer and/or donor complaints?			Ranking





Н-	- Release Procedures			
#	Audit Question	Yes	No	Ranking
1.	Does QA/responsible person have the authority and			Critical
	responsibility as outlined by the plasma center SOP or			
	job description to stop			
	a) the release of plasma for shipment, if necessary?			
-	b) plasma center production, if necessary?			Oritical
2.	Does the company have written procedures that outline			Critical
	and instruct QA/responsible person on the specific checks that must be verified as acceptable before			
	plasma units are released?			
3.	Is final plasma release controlled by Quality Assurance			Critical
0.	personnel or a qualified alternate?			ondoar
Au	ditor Comments on Section H:			
1-	Viral Marker Standard			
- #	Viral Marker Standard Audit Question	Yes	Νο	Ranking
	Audit Question Does the company have written procedures in place	Yes	No	Ranking Critical
# 1.	Audit Question Does the company have written procedures in place that conform to the IQPP Viral Marker Standard?	Yes	No	Critical
#	Audit QuestionDoes the company have written procedures in place that conform to the IQPP Viral Marker Standard?If the data are computer-generated, is the query set up	Yes	No	
# 1.	Audit QuestionDoes the company have written procedures in place that conform to the IQPP Viral Marker Standard?If the data are computer-generated, is the query set up to classify Applicant and Qualified Donors according to	Yes	No	Critical
# 1. 2.	Audit QuestionDoes the company have written procedures in place that conform to the IQPP Viral Marker Standard?If the data are computer-generated, is the query set up to classify Applicant and Qualified Donors according to the IQPP Qualified Donor Standard?	Yes	No	Critical
# 1. 2.	Audit QuestionDoes the company have written procedures in place that conform to the IQPP Viral Marker Standard?If the data are computer-generated, is the query set up to classify Applicant and Qualified Donors according to	Yes	No	Critical
# 1. 2.	Audit QuestionDoes the company have written procedures in place that conform to the IQPP Viral Marker Standard?If the data are computer-generated, is the query set up to classify Applicant and Qualified Donors according to the IQPP Qualified Donor Standard?	Yes	No	Critical
# 1. 2.	Audit QuestionDoes the company have written procedures in place that conform to the IQPP Viral Marker Standard?If the data are computer-generated, is the query set up to classify Applicant and Qualified Donors according to the IQPP Qualified Donor Standard?	Yes	No	Critical
# 1. 2.	Audit QuestionDoes the company have written procedures in place that conform to the IQPP Viral Marker Standard?If the data are computer-generated, is the query set up to classify Applicant and Qualified Donors according to the IQPP Qualified Donor Standard?	Yes	No	Critical
# 1. 2.	Audit QuestionDoes the company have written procedures in place that conform to the IQPP Viral Marker Standard?If the data are computer-generated, is the query set up to classify Applicant and Qualified Donors according to the IQPP Qualified Donor Standard?	Yes	No	Critical
# 1. 2.	Audit QuestionDoes the company have written procedures in place that conform to the IQPP Viral Marker Standard?If the data are computer-generated, is the query set up to classify Applicant and Qualified Donors according to the IQPP Qualified Donor Standard?	Yes	No	Critical
# 1. 2.	Audit QuestionDoes the company have written procedures in place that conform to the IQPP Viral Marker Standard?If the data are computer-generated, is the query set up to classify Applicant and Qualified Donors according to the IQPP Qualified Donor Standard?	Yes	No	Critical
# 1. 2.	Audit QuestionDoes the company have written procedures in place that conform to the IQPP Viral Marker Standard?If the data are computer-generated, is the query set up to classify Applicant and Qualified Donors according to the IQPP Qualified Donor Standard?	Yes	No	Critical
# 1. 2.	Audit QuestionDoes the company have written procedures in place that conform to the IQPP Viral Marker Standard?If the data are computer-generated, is the query set up to classify Applicant and Qualified Donors according to the IQPP Qualified Donor Standard?	Yes	No	Critical
# 1. 2.	Audit QuestionDoes the company have written procedures in place that conform to the IQPP Viral Marker Standard?If the data are computer-generated, is the query set up to classify Applicant and Qualified Donors according to the IQPP Qualified Donor Standard?	Yes	No	Critical





J –	Cross Donation Management Standard			
#	Audit Question	Yes	No	Ranking
1.	Does the company have written procedures that conform to the IQPP Cross Donation Management Standard and contain the following?			Critical
	a) A procedure stating how it will prevent an individual found to be an active donor in more than one center from donating more often than allowed.			
	b) A notification process to inform all known plasma centers within a center's DRA of the opening of a new center and provide all required information to the CDCS no later than 30 days prior to the scheduled opening date.			
	 An articulated process to apply a permanent deferral to a donor who is found to be cross- donating. 			
	d) SOP requiring use of the PPTA Cross Donation Check System (or, where the CDCS is not permissible by law, an alternative national or regional registry, if available, and, where no alternate deferral registry is available, an intra- company process) in accordance with the Standard.			
<u>Auc</u>	litor Comments on Section J:			



K -	Standard for Recording Donor Adverse Events			
#	Audit Question	Yes	No	Ranking
1.	Does the company have a documented process for			Major
	recording known DAEs considered to be associated			2
	with any part of a Source Plasma donation program			
	(this includes initial screening, donation, immunization			
	for high titer collections, etc.) following company			
	approved SOPs, and does this process conform to the			
	Standard?			
2.	Does the process require centers to record, in the			Major
	facility's documentation system, DAEs as required by			
	the Standard?			
Au	ditor Comments on Section K:			





General Overall Comments:

Ranking Guidelines:

Critical Observations	= 50 points each
Major Observations	= 10 points each

Scoring Guidelines:

A score of 51 points or more will trigger a procedure in which a re-audit in less than one year may occur.

