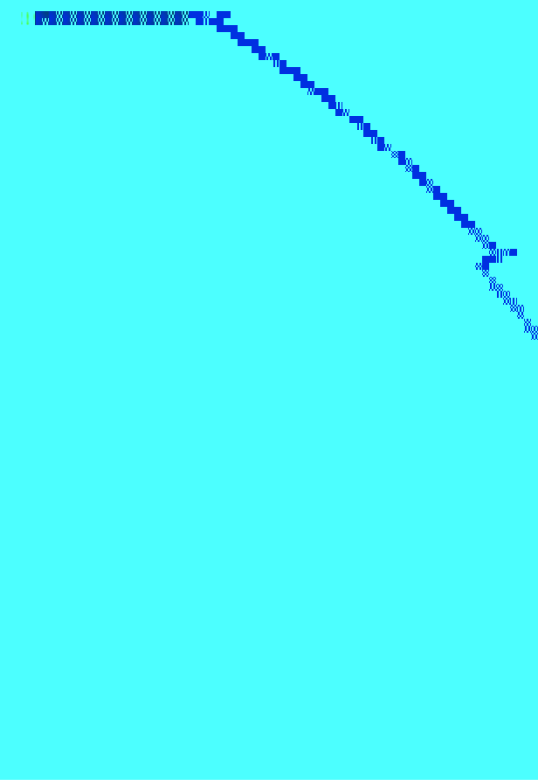


Non-Hospital Medical and Surgical Facilities Accreditation Program

ACCREDITATION STANDARDS

Allografts



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No.	Description
ALO1.0	ALLOGRAFTS
A	

No.	Description
ALO1.4.2	<p>M There is policy and procedures for the investigation and reporting of incidents involving allografts. <i>Guidance: The policy and procedures outline the investigation of complications, technical problems, errors, accidents or adverse reactions involving allografts, the reporting of complications, technical problems, errors, accidents or adverse reaction involving allografts to the source establishment or distributor and any corrective action taken. Allograft incidents that reach the patient, both no harm and harm events, are also reported to the College in accordance with the bylaws for patient safety incidents. Investigation and reporting of incidents involving allografts are maintained for, at minimum, 16 years.</i></p>
ALO1.4.3	<p>M There is policy and procedures for lookback notifications and recall of allografts. <i>Guidance: The policy and procedures outline the regulated health professional(s) responsible for lookback and recall activities, acknowledgement of lookback or recall notification, notification of the physician of the patient or the patient directly if required, and quarantining of allografts in inventory until final disposition is determined. A lookback is the tracing and testing of allograft recipients in cases where the allograft is determined to be potentially contaminated with a blood-borne infection. A recall is a notification by the source establishment or distributor when a quality problem requiring action has been identified. Lookback notification and recall records are maintained for, at minimum, 16 years.</i></p>

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Revision history

Date	Revisions
July 4, 2019	The following changes were made: