

# New EU requirements for MUDI Updates MSWG

Project Lead: Greg Rowe

OGSM  
Report Date: December 2025  
Expected project close date Q2 2026

## Business purpose

- This group will develop the appropriate solution. The European (EU) Commission submitted a work request to develop the creation of "Master UDI-DI" for spectacle frames, spectacle lenses and ready-to-wear reading spectacles as part of the UDI requirements based on the European Medical Device Regulation (MDR).

## Key issues and risks

- EU COM delegated act has not been finalised. The team is moving forward with the current understanding of the potential language in the delegated act which could impact the draft GSCN.

## Accomplishments this period

- Team has been working on the conducting the Community Review Comment Resolution process
- The team is expected to complete the process on the 09 December team call and motion the BRAD to eBallot

## Upcoming period activities

- Motion BRAD to eBallot
- Begin working on the Solution document

# New EU requirements for medical devices identification MSWG

## Overall Status

#	Standard /guideline Milestones	Due Date	Notes	#	Collateral milestones	Due date	Notes
1	Master UDI-DI GSCN	Q2 2026		1	TBD	TBD	
2							
3							
4							
5							

### Stakeholders

### Project Scope, Resources and Timeline Change Management

Sponsor	Robert Beideman			Kick-off call May 2025					
SDL	Greg Rowe								
CE/Sol Liaison									
SME	Neil Piper								
AG Liaison	Shekhar Nambi – J&J								
Chairs	Lionel Tussau – Bayard Consulting, Sylvia Reingardt – GS1 Germany, Jeremie Consoln								
New risks identified	Yes/no	Change request included	Yes/no	Legend	Complete	Not Started	On target	Risk of being late	Change needed (not recoverable)