

Visual Inspection of Injectable Products:

Myth Busting ...

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- Inspection Myths
- Conclusions
- References and Acknowledgements



Inspection Myths



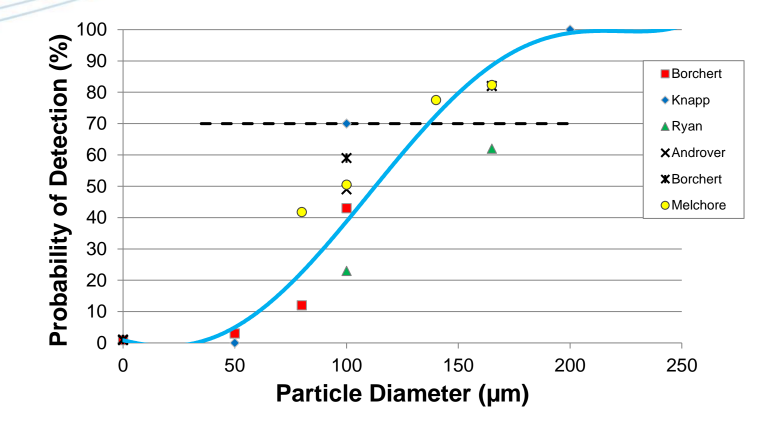
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PDA Inspection Myth #1

- 100% inspection means detection and elimination of all visible defects (e.g. particulate matter, cracks, etc.)
 - Inspection is a probabilistic process.
 - Detection probability is dependent on inspection conditions and defect characteristics.
 - Particles <200 um generally have a detection probability <100%.



Human Inspection Performance



From Shabushnig, Melchore, Geiger, Chrai and Gerger, PDA Annual Meeting 1995

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PDA Inspection Myth #1

- 100% inspection means detection and elimination of all visible defecting. particulate matter, cracks, tc.)
 - Inspection is a probability of the set
 - Detection probable desensant on inspection condition, an defect o aracteristics.
 - Particles 22 Concenerally have a detection probability 200%.

Inspection Myth #2

- Human manual inspection is a "validatable" process.
 - Human inspectors are not
 - Qualified human implate call in vide reliable performance
 - Definerts econ and trailing criteria
 - Control a spl tion conditions
 - Light ckground, Duration
 - SOP's

Inspection Myth #3

- Magnification always improves human manual inspection performance.
 - Inspectors will move head position to minimize eyestrain during extended inspection, reducing apparent magnification.
 - Controlled studies have not found increased detection of particulates or container defects with 3x magnification.



Detection Rate with Magnification

	5 mL		30 mL	
	No Mag	Mag	No Mag	Mag
Product	50.0%	37.5%	18.6%	18.6%
Container	37.5%	37.2%	45.4%	44.6%
Closure	62.3%	54.2%	72.5%	68.2%
All Defects	50.6%	46.0%	53.6%	51.4%
Good	0.5%	0.9%	2.0%	0.6%

Semi-automated inspection at 55 VPM, lyo test set, n=1000, 3x mag

Inspection Myth #3

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- If you use a sampling plan with an AQL of 0.1% and do not exceed the acception per in your sample, the defect rate is your by convill not exceed 0.1%.
 - AQL is the Acceptable value Level and is the defect rate where the ejection probability is 5%. 95% of batches where the belieft rate will be <u>accepted</u>. This is a measure for erisk of rejecting good batches.
 - The UQL is the Unacceptable Quality Level and is the defect rate where the rejection probability is 90% for the batch.



Conclusions

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13



- Current industry performance is generally at or beyond the limits of medical risk.
- Compendial guidance is ambiguous, but getting better.
- "Zero defects" is a valuable goal, not a practical limit for particulate matter.
- Need to develop practical limits based on risk assessment and process capability measures.



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 - Subpart D Equipment
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 - 211.94 Drug product containers and closures
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 - 211.110 Sampling and testing of in-process materials and drug products
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Conferences and Meetings

- PDA Visual Inspection of Parenterals Interest Group
- PDA Visual Inspection Forums

Equipment Vendors

- **Antares Vision**
 - Brescia, Italy
- Brevetti C.E.A., S.p.A.
 - Sovizzo, Italy www.brevetti-cea.com
- Bonfiglioli Engineering, S.r.l.
 - Vigarano Pieve, Italy www.bonfiglioliengineering.com
- Dabrico, Inc.
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- eyetec
 - Antwerp, Belgium www.eyetec.be
- Syntegon Technology, GmbH (formerly Eisai, Bosch)
 - Waiblingen, Germany <u>www.syntegon.com</u>
- InnoScan K/S (Stevenato Group)
 - Braband, Denmark
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www.antaresvision.com

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 - Padova, Italy www.optrelinspection.com
- Phoenix Imaging

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- Livonia, MI
 www.phoeniximaging.com
- Seidenader, GmbH (Korber)
 - Munich, Germany www.seidenader.de
- Unchained Labs (Rap.ID Particle Systems)
 - Pleasanton, CA www.unchainedlabs.com
- Lighthouse Instruments
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www.lighthouseinstruments.com

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 - Warrington, PA

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- Micro Measurement Laboratories, Inc.
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- Prime Results
 - Harrisburg, PA

www.phoeniximaging.com

www.prime-results.com

- SoloHill Engineering, Inc
 - Ann Arbor, MI

www.particlestandards.com

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Questions



Remember, everyone is an inspector ...

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