



Mastering AVI

Part 5: transition from MVI to AVI



Instructor Lead: Romain Veillon / Fernand Koert / Sébastien Koch

© Copyright PDA Author Romain Veillon



MVI remains Golden Standard

« The reference method described in this chapter and in <790> is a manual inspection of a single container for particulate matter “



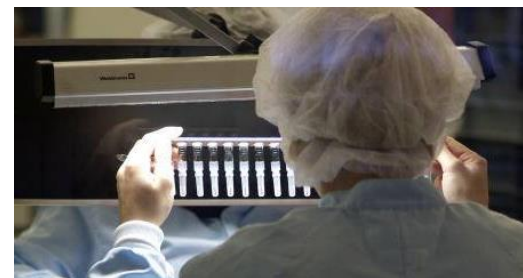
“However, multiple-container manual inspection, semi-automated, and automated inspection methods are also discussed and permitted by the Pharmacopeia. These alternate inspection methods must be qualified to demonstrate equivalent or better defect detection when compared to the reference manual inspection described in <790>.”



“Holding many containers by hand at once should be avoided, as it is difficult to obtain a complete view of all container surfaces and contents. Full rotation (360°) of the container during the container–closure defect inspection sequence is recommended for identifying small container defects such as cracks or chips”



“
If multiple containers are qualified to be equivalent to the single-container inspection method per <790>, they may be held during the particle detection sequence using a tool that holds these containers for consistent presentation.”



“Qualification of inspectors and validation of the inspection equipment should be based on comparison with the compendial single-container manual-inspection process with an expectation that alternative methods such as semi-automated inspection demonstrate equivalent or better performance”





Automated Visual Inspection (AVI)

- ✓ High speed and high capability
 - ✓ Highly reproducible
 - ✓ Consistent (no fatigue effect)
 - ✓ Defects presentation
-
- ✓ High initial investment
 - ✓ Works within strict condition (validated upstream process)
 - ✓ Indiscriminative (i.e.: fiber and cracks are seen the same way)
 - ✓ Some uncovered area
 - ✓ Higher false reject rate
 - ✓ No classification



Semi-Automated Visual Inspection (SAVI)

- ✓ Adaptation
 - ✓ Speed
 - ✓ Brain
 - ✓ Flexible
 - ✓ Decision capable
-
- ✓ Inconsistent (fatigue effect)
 - ✓ Not highly reproducible
 - ✓ Susceptible to influence
 - ✓ Some uncovered area
 - ✓ Monotonous repeated work
 - ✓ Significant training effort

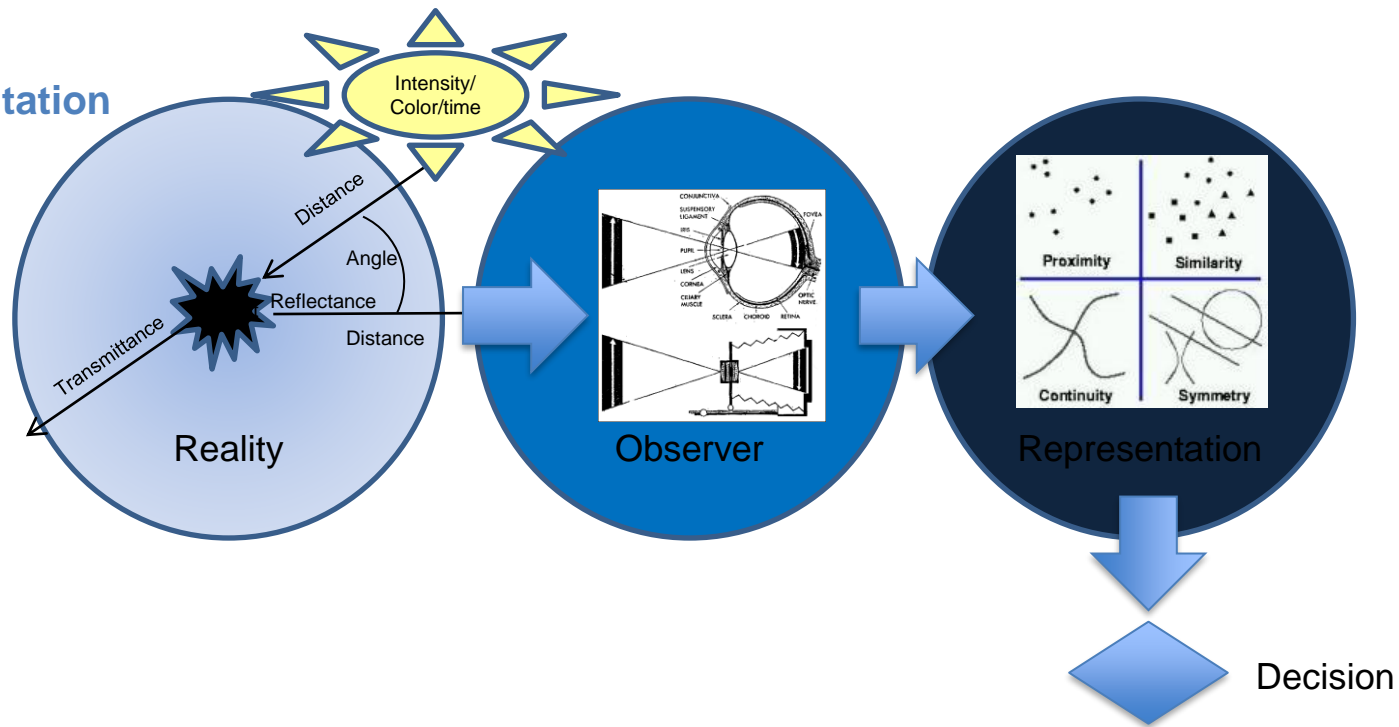


Manual Visual Inspection (MVI)

- ✓ Adaptation
 - ✓ Brain
 - ✓ Flexible
 - ✓ Decision capable
 - ✓ Classification of defects
-
- ✓ Inconsistent (fatigue effect, emotional)
 - ✓ Not highly reproducible
 - ✓ Susceptible to influence
 - ✓ Slow
 - ✓ Monotonous repeated work

Inspection steps from object presentation to decision

Object presentation



Example of MVI interpretation with color continuity : SNOW can be blue ?

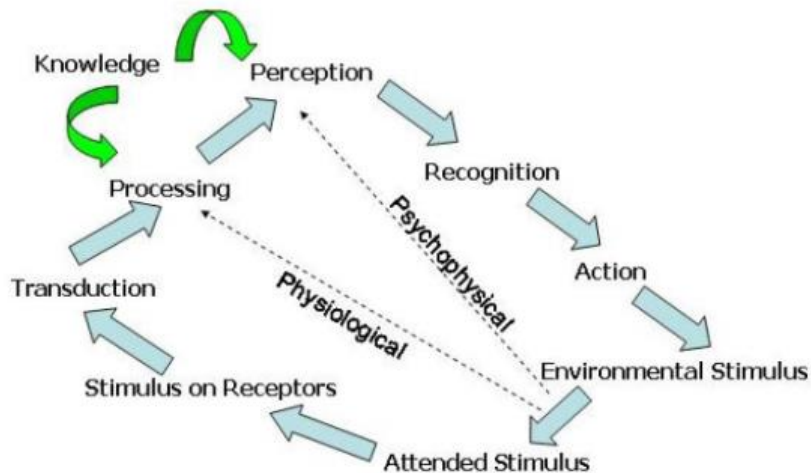
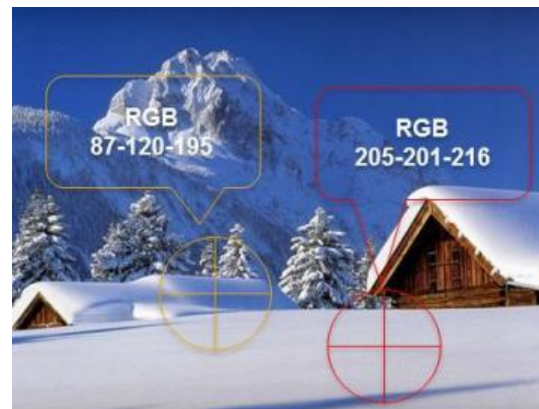
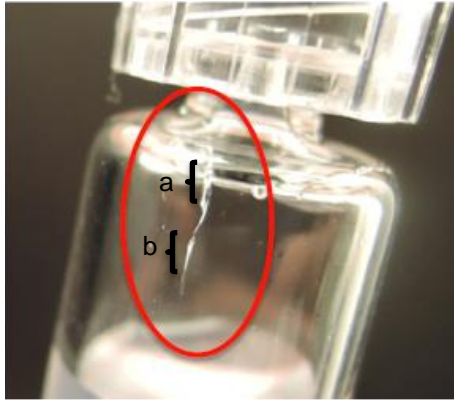


Figure 1.1: The Perceptual Process

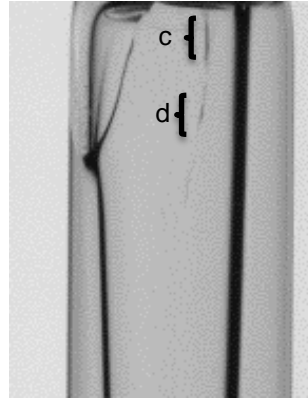


Chromatic continuity:
We see snow even when color
changes drastically (RGB)

Example of MVI interpretation of a crack versus machine



The inspector will be attracted due to the glistering of a and b. The human brain will, out of experience, connect a and b over the thin lines making it a medium crack



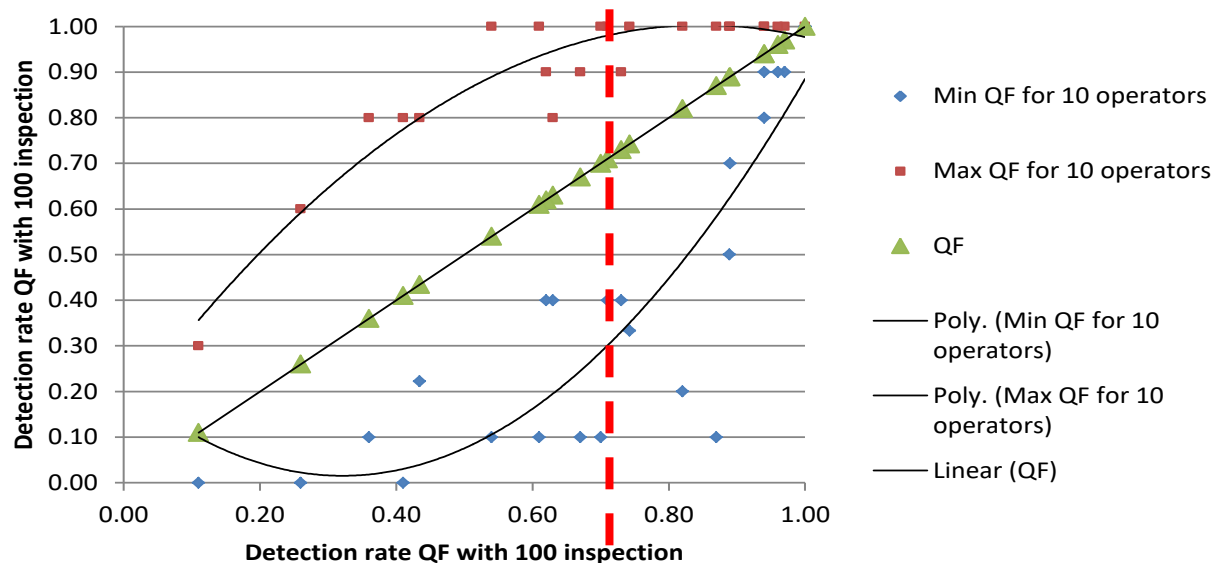
A crack becomes visible when the crack is under a certain angle towards the camera because light cannot pass the crack, creating a shade, c and d.

The thin lines are hardly or not visible at all, so c and d are not connected making it two minor cracks. One can push it but this will automatically imply that small scratches are also rejected.

C and d can be detected without, but when comparing human and machine one compares medium with minor.

MVI inter-operator variability increases with smaller defects

QF distribution for Syr. Particle defects



Key learning:
 Particle detection in MVI is highly probabilistic: operator variability is lower with very high QF > 0.70
 Operator variability higher with lower QF [0.3:0.8]
 To compare AVI to MVI need to be in true defect zone where inter-operator variability is lower

Establish MVI baseline

Manual Visual inspection Baseline study (Knapp)

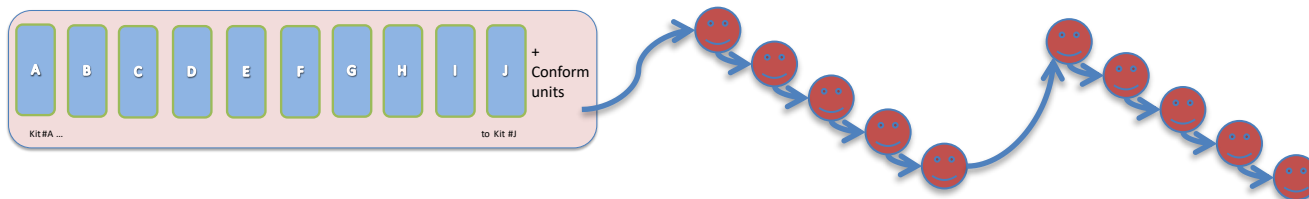


Example of standard MVI Baseline evaluation

Key learning:
Proposal for methodology for
MVI baseline evaluation

Material and Methods

- [10 kits + good units] = 1 inspection lot order
- No information given to inspectors = routine inspection
- No interactions with inspectors to avoid any interferences
- Changed shift to avoid interactions between inspectors
- 1 inspection every day during 2/3 weeks, one inspector at a time
- Kit verified every day for defect state, replaced broken units to identical
- QF Result compiled for each inspector















Example of standard MVI Baseline evaluation

Material and Methods

Data reporting: QF = number of ejected / number of inspected

	Operators										
		1	2	3	4	5	6	7	8	9	10
KIT	DEFECT										
Kit A	Defect #1										
Kit A	Defect #...										
Kit A	Defect #nn										
.....										
Kit J	Defect #1										
Kit J	Defect #...										
Kit J	Defect #nn										

QF #1A

.....
QF #nn

QF #1J

QF #nn

MRZE

Key learning:
 Rigorous Baseline evaluation of MVI performance is key to succeed AVI validations
 Mind Data integrity control
 Good documentation practices

Knapp Digested

Since the particulate visibility statement in the XIX Revision of the Pharmacopeia (9) is based upon a deterministic human inspection it is inappropriate and should be discarded.

With both manual and automated systems regarded as probabilistic, they can now be similarly evaluated and their demonstrated capability rigorously compared.

longevity estimates. The particular containers rejected in any single inspection cannot be accurately predicted except for two special cases: those containers that are absolutely clean and are never rejected and those containers with gross defects that are rejected in every inspection.

Key learning:
MVI and AVI remain
probabilistic by nature

Uhlir pioneer work for method comparison: Venn diagram

In terms of the two-dimensional probabilistic inspection model, Uhlir utilized two unrelated one-dimensional probability distributions: manual and machine. In consequence, the differing sensitivities of the two methods can yield the Venn diagram result shown in Figure 1. Here, the manual inspection and the automated device perform in exemplary fashion. Figure 1 indicates, however, that the sets of containers rejected by each method had few containers in common. This comparison suggests that the Uhlir evaluation methodology may not generate the demonstration of equivalence that CGMP's require in the validation of alternative inspection methodologies and devices.

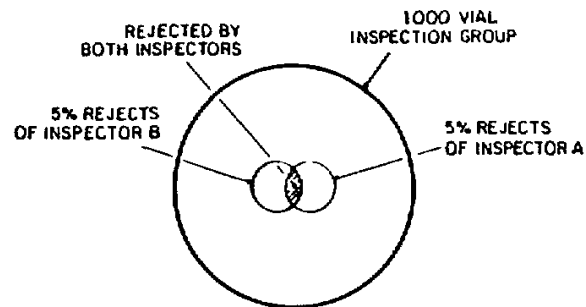
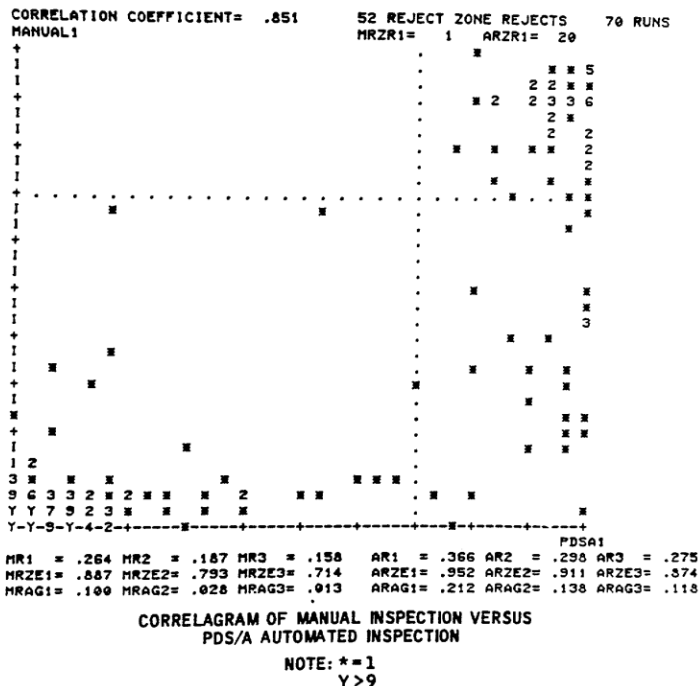


Figure 1—Venn diagram of two inspector particulate inspection demonstrating the expected paradoxical results.

Key learning:

Between multiple operators in MVI there is no contingency, meaning even if they have similar performance, they cannot not detect same defects

Why Correlogram unit by units does not make sense ?



Key take Away:
 When J Knapp draw a correlogram of between 2 method, each plot is the number of units in each probability class
That is NOT paired comparison per DEFECT

The capability of one process relative to the other cannot be evaluated until the correlation between the results of both inspections is established. This correlation is based on an examination of the inspection history of each container in each inspection process. Sufficient inspection replications are required to assure statistically reproducible results with acceptable tolerance intervals. Since we are dealing with probabilistically defined quantities, statistical tools must be used. The basic questions of replicability, relative per-

Figure 3—Correlogram comparing the results of 72 manual and 70 PDA/A inspections. A comparison summary of the two inspection methods is included in the computer printout. Of major interest is the fact that only 1 (MRZR1) of the 52 were rejected manually with a probability of 1.0. The PDA/A rejected 20 (ARZR1) of the 52 with a probability of 1.0. The plusses on each axis are the 10% increment points from a rejection probability of 0 to 1.0. The abscissa is for the automated system; the manual system rejection probability is the ordinate. The dotted lines shown are the Reject Zone boundaries for both systems. The * symbols indicate a single container at a point in the plane, a Y indicates a number of containers greater than 9. Values between 2 and 9 are shown directly. The reject rate, R, the Reject Zone Efficiency, RZE, and the undesired reject rate in the Accept and Gray Zones, RAG, are tabulated under the histogram with suffix 1, 2, and 3 to indicate sequential inspection number. The prefix N indicates manual inspection; the A prefix indicates an automatic inspection.

Classification of defects by « iso-probability subgroup »

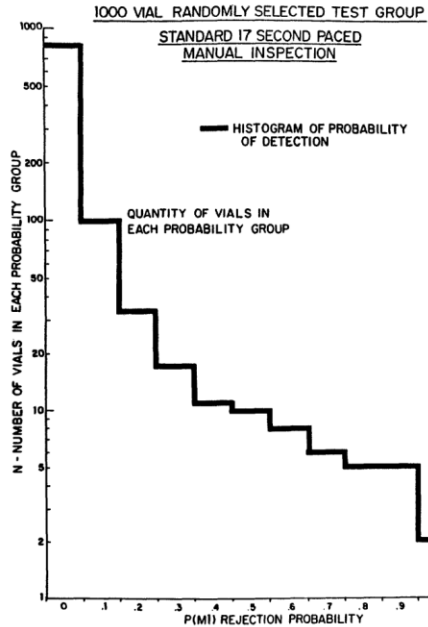


Figure 2—Histogram of probability of detection for a 1000 vial randomly selected test group. The Schering standard 17 second paced manual inspection was employed.

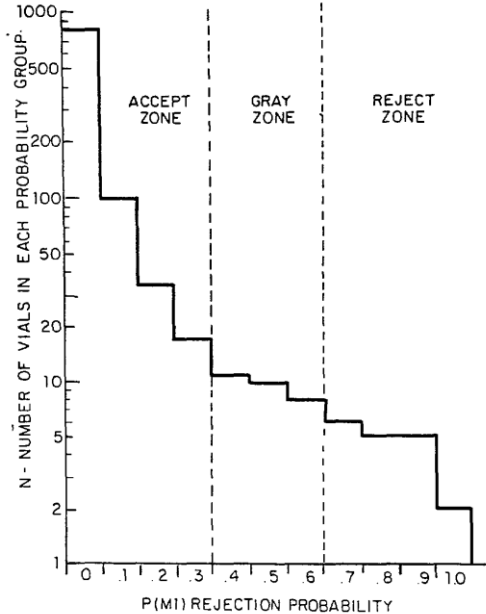


Figure A2—One-dimensional histogram of a normal batch showing the accept Gray and Reject Zones defined by the human based standard inspection.

How to compare 2 distribution of probability?

manual inspection capability. To accomplish this evaluation, two random distributions must be compared.

When the implications of the two dimensional probability plane of Figure A2 are examined it becomes apparent that each entry in either system can be transformed into a distribution in the other system.

TABLE AII. Probabalistic Distribution of Rejection Probabilities for Containers in "Manual" Inspection and "System" I and II

	N(0)	N(.1)	N(.2)	N(.3)	N(.4)	N(.5)	N(.6)	N(.7)	N(.8)	N(.9)	N(1.0)
"MANUAL"	1	1	1	1	1	1	1	1	1	1	1
"SYSTEM"	2.5	1.5	1.5	.5	0	0	0	.5	.5	1.5	2.5
I & II											

RZEM terminology

$$RZE(M1) = \frac{RZR(M1)}{RZN} = \frac{14.7}{18} = 81.7\% \quad (\text{Eq. 4})$$

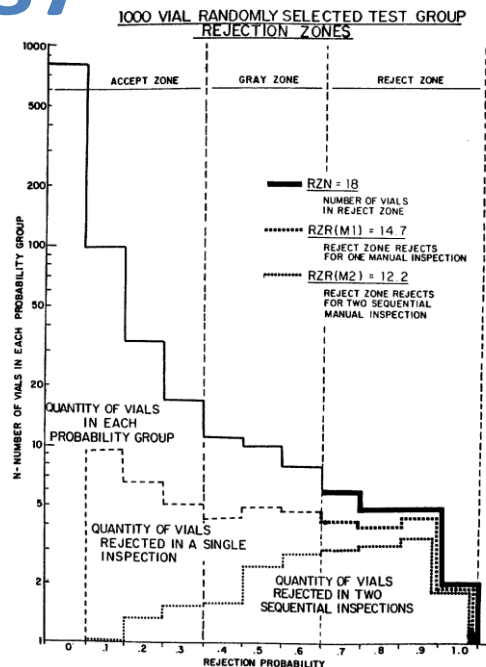
$RZE(Mn)$ = efficiency of rejection in Reject Zone

$RZN(Mn)$ = number of vials identified in the manual Reject Zone

$RZR(Mn)$ = Reject Zone reject quantity as defined in manual inspection

$$RZE(M2) = \frac{RZR(M2)}{RZN} = \frac{12.2}{18} = 67.7\%$$

RZE = Reject zone efficiency



Key take Away:
Now USP has simplified terminology speaking of PoD Probability of Detection

AVI validation approaches

- Comparison AVI to MVI baseline: Knapp approach
- AVI to be better or equivalent to MVI pre established baseline on true defects

- Fixed criteria for AVI validation

i.e

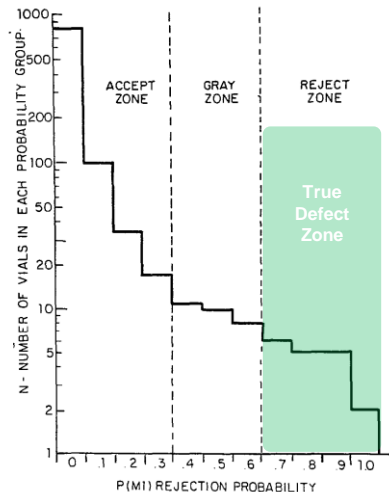
- critical > 90%
- Major >70%
- FRR < 5%

- Fixed criteria is not meaning 100% !

Comparison AVI to MVI

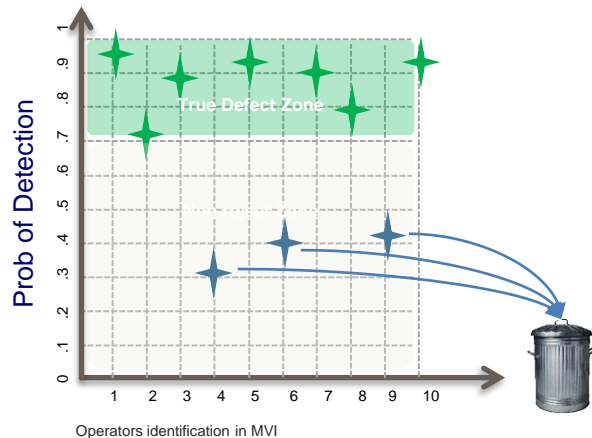
Knapp concept of true defect zone

- ❑ Performance of a new method (MVI – SAVI – AVI) must be compared to MVI Baseline PoD established with standard work conditions
- ❑ Only defects in the true defect zone are retained, when PoD is above 70%
- ❑ Comparison is not a paired comparison defect by defect but rather based on average comparison for a defect family (number of vials in each probability group)

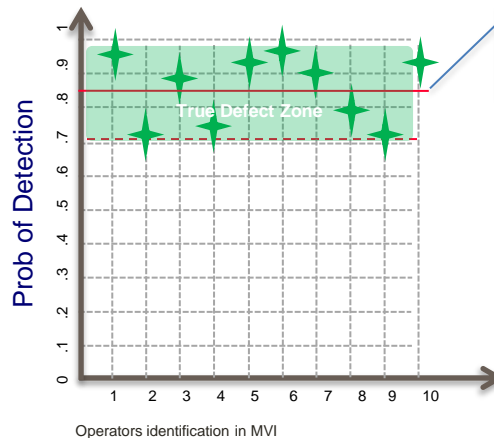


True Defect Zone concept (USP<1790> Knapp)

MVI Baseline performance study by defect family



MVI Baseline performance study by defect family
With only true defect zone (Pod>0,7)

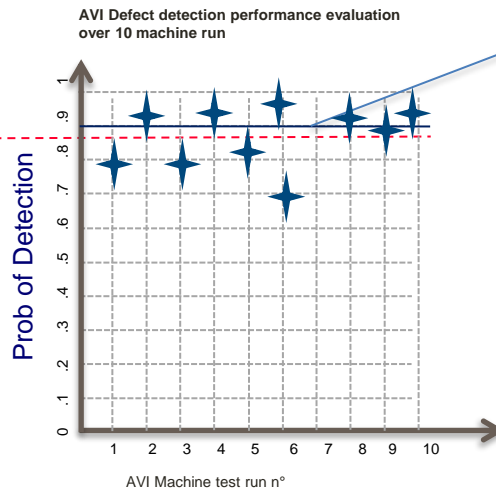
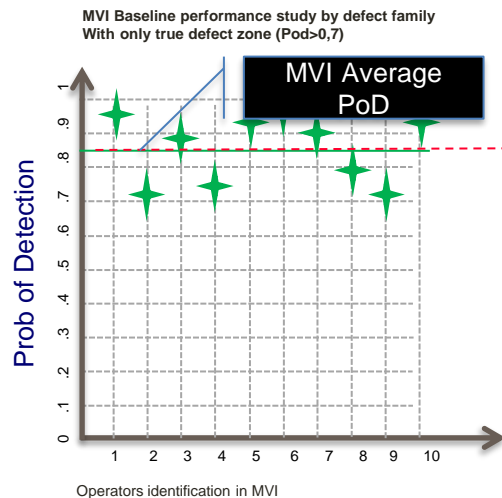


Average PoD
MVI

Critical Design Element:
Recent FDA guidance on particle insist that when particle are below 70%, the MVI st Work should be challenged rather than just not considering the units

Comparison AVI to MVI

“The capability of one process relative to the other cannot be evaluated until a correlation between the results of both inspections is established. This correlation is based on an examination of the inspection history of each container in each inspection process. Sufficient inspection replications are required to assure statistically reproducible results with acceptable tolerance intervals. Since we are dealing with probabilistically defined quantities, statistical tools must be used.” J. Knapp

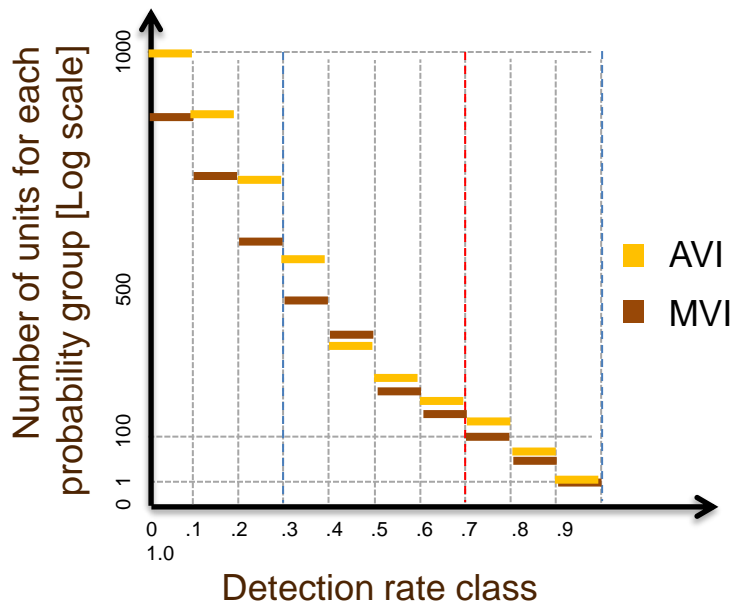


AVI average PoD is compared to MVI baseline

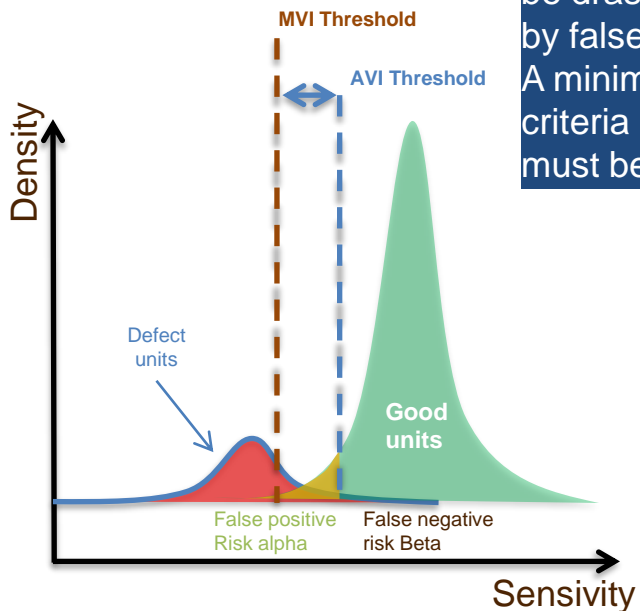
Some Parameters for PQ Design

Why it is critical to control false reject?

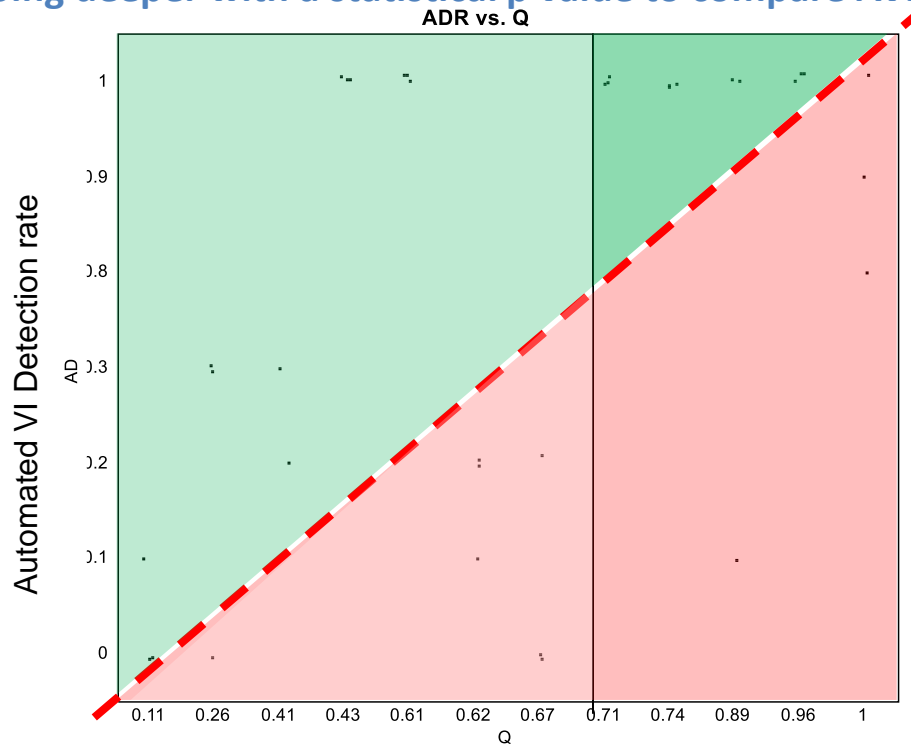
Key take Away:
PoD of defect should not be drastically increased by false reject.
A minimum validity criteria of false reject must be established



=> Comparison of 2 distributions of number of unit having same detection rate



Going deeper with a statistical p value to compare AVI vs MVI



Key learning:

Particle detection in AVI has a higher ADR and is less probabilistic than MVI
Specially in range of QF > 0.70

In range with Lower QF
ADR is higher than MVI but more heterogeneity between particles (floating/precipitating)

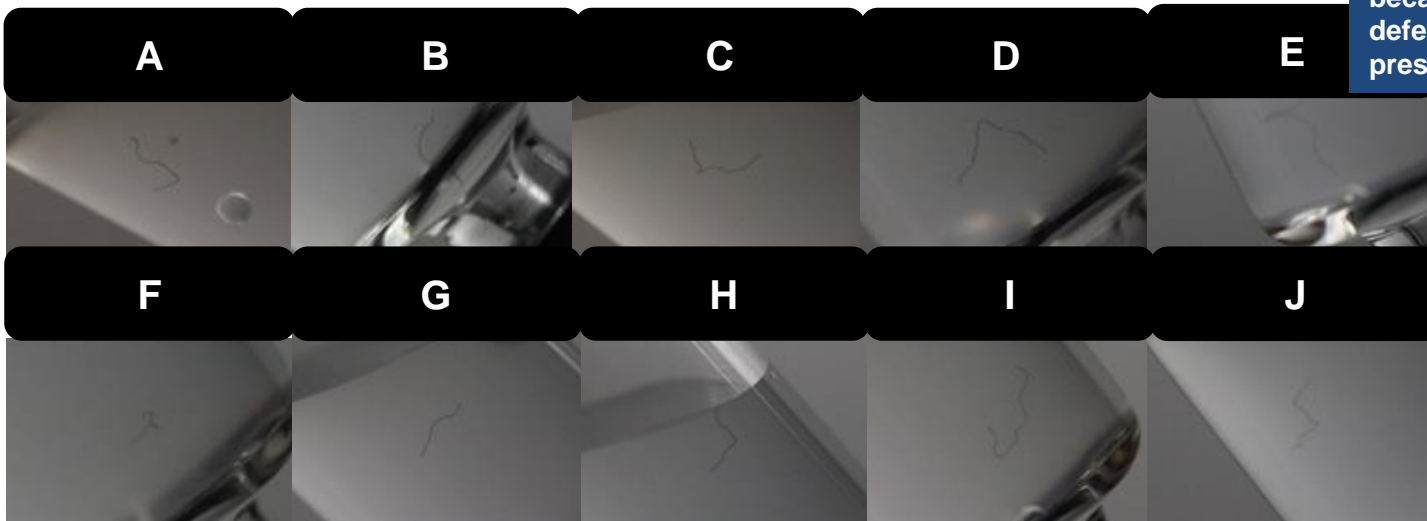
Knapp demonstrated that Validation comparison AVI to MVI should be done in True Defect Zone using “gross defects”

Some individual defect may be lower in detection on some run, the average probability of defect for a defect type (ie particle) must be considered rather than individual paired comparison defect by defect.

Replicate

"The availability of an adequate number of vials in each rejection probability set will be seen to be a prerequisite for successful validation experiments." J.Knapp

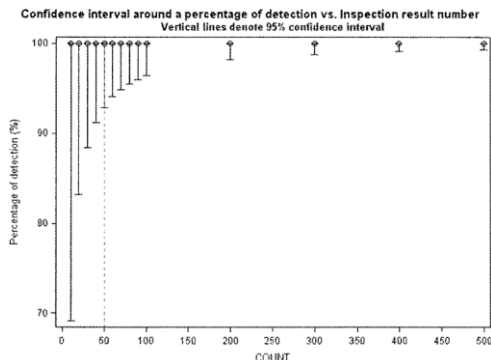
Key learning: At least 3 replicates per defect type should be considered for validation because variability of defect + defect presentation



Impact of number of validation run

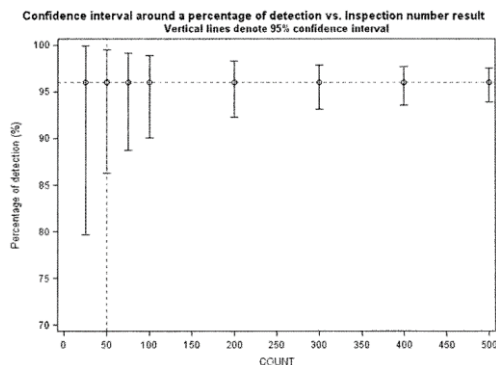
Sample size: practical impact in test run design

With Detection rate limit 100%



With hypothesis of binomial distributions
With 50 runs in validation the confidence interval at 95% is: [92.9% ; 100%]

With Detection rate limit 96%



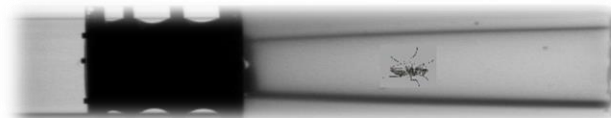
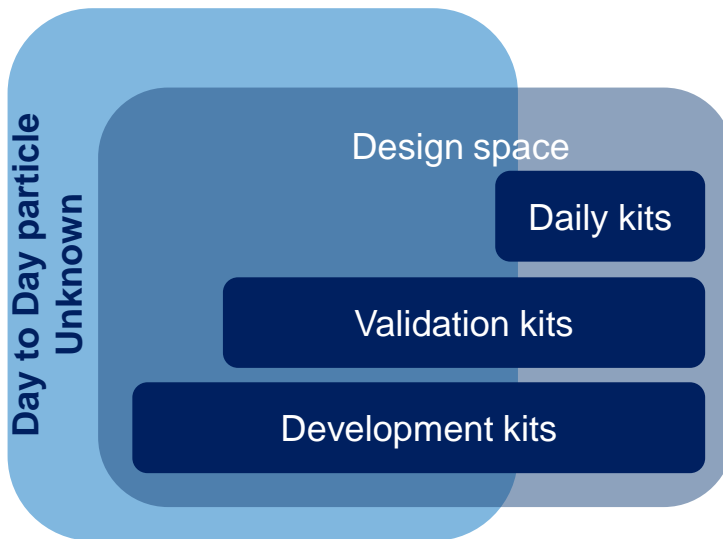
With hypothesis of binomial distributions
With 50 runs in validation the confidence interval at 95% is: [83.6% ; 99.5%]

Key learning: even in case of non probabilistic detection rate criteria, the result remains in a Conf. Int. that depends of number of validation runs

Ability for unknown defects



- Machine vision is designed with minimum threshold, may be compared to high jump.
- Machine vision is designed to detect defects that are outside the design space to anticipate some new defects (unknown)
- With artificial image library we can demonstrate capability of unknown detection (I.e extrinsic)



!example = Fake image!

RECAP



You have learnt

AVI vs MVI

- Machine qualification
- Interpretation of inspection results and validation data : Knapp review
- Considerations on validation program for automated inspection
- Performance measurement
- Maintaining the manual inspection

AVI vs MVI

- How many operators do you need for a baseline study
- How many inspections for a subtype is minimal needed
- What should be the minimal detection rate for MVI-AVI comparison
- Which zones had Knapp defined and which one should we avoid
- How do we compare MVI-AVI