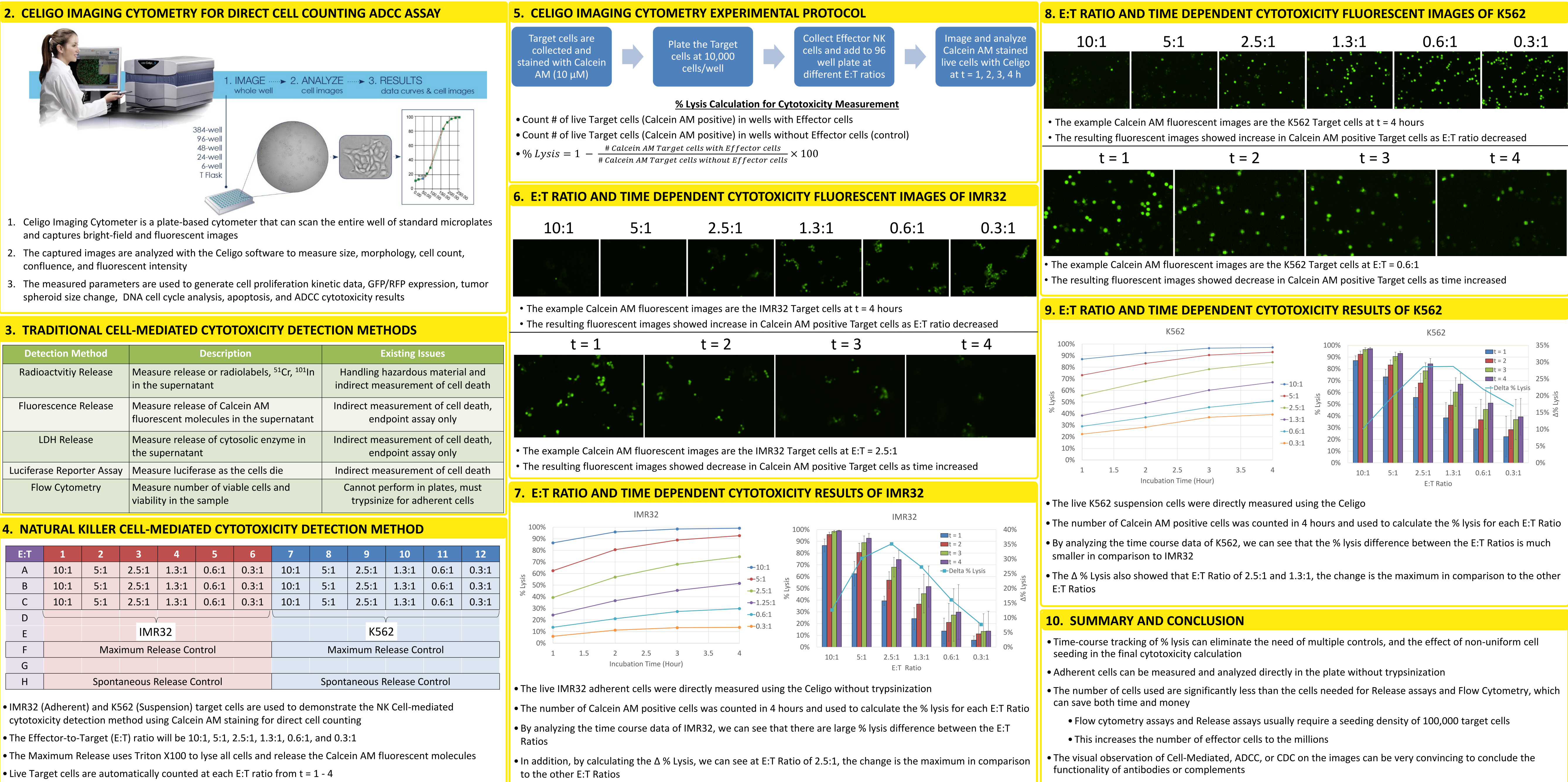
# **Revolution**

### **1. ABSTRACT**

Cytotoxicity assays play a central role in studying the function of immune effector cells such as cytolytic T lymphocytes (CTL) and calcein release assays. The assays involve labeling tumor cells (target) with radioisotope or fluorescent dyes, when the target cells are subjected to cytolysis by CTLs or NK cells (effector), they release the entrapped labels into the media upon lysis. The amount of labels in the media is measured to determine the level of cytotoxicity the effectors have induced. poor loading efficiency and high spontaneous release of the reagents. In this work, we demonstrate a novel cytotoxicity assay using the Celigo imaging cytometry, direct cell counting of live fluorescent target cells can be performed, which is a direct method for assessment of cytotoxicity. Human NK cells from one healthy donor were used as effectors, and K562 (suspension) and IMR32 (adherent) were then added to each well at Effector-to-Target (E:T) ratios 10:1, 5:1, 2.5:1, 1.25:1, 0.625:1, and 0.3125:1. The 96 well plate was then scanned and analyzed using Celigo imaging cytometer at t = 1, 2, 3, and 4 h to measure the % lysis of target cells. The results showed increased. The proposed Celigo imaging cytometer at t = 1, 2, 3, and 4 h to measure the % lysis of target cells. cytotoxicity, which can be an attractive method for both academic and clinical research.



- and captures bright-field and fluorescent images
- confluence, and fluorescent intensity

## **Detection Method**

Radioactvitiy Release	Measure release or radiolabels, <sup>51</sup> Cr, <sup>101</sup> In in the supernatant	Handling haz indirect meas	
Fluorescence Release	Measure release of Calcein AM fluorescent molecules in the supernatant	Indirect meas endpo	
LDH Release	Measure release of cytosolic enzyme in the supernatant	Indirect meas endpo	
Luciferase Reporter Assay	Measure luciferase as the cells die	Indirect meas	
Flow Cytometry	Measure number of viable cells and viability in the sample	Cannot per trypsinize	

	E:T	1	2	3	4	5	6	7	8	9	
	А	10:1	5:1	2.5:1	1.3:1	0.6:1	0.3:1	10:1	5:1	2.5:1	
	В	10:1	5:1	2.5:1	1.3:1	0.6:1	0.3:1	10:1	5:1	2.5:1	
	С	10:1	5:1	2.5:1	1.3:1	0.6:1	0.3:1	10:1	5:1	2.5:1	
Ĩ	D				(		)				
	E			IMF	32					K56	52
	F	Maximum Release Control						Maximum Re			
	G										
	Н		Spont	aneous F	Spontaneous Re						

cytotoxicity detection method using Calcein AM staining for direct cell counting

- The Effector-to-Target (E:T) ratio will be 10:1, 5:1, 2.5:1, 1.3:1, 0.6:1, and 0.3:1
- Live Target cells are automatically counted at each E:T ratio from t = 1 4

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## Quantification of Natural Killer Cell-Mediated Cytotoxicity using Celigo Imaging Cytometry

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