

Quantification of Colibactin-associated Genotoxicity in HeLa Cells by In Cell Western (ICW) Using γ -H2AX as a Marker

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[Abstract] The genotoxin colibactin is produced by several species of *Enterobacteriaceae*. This genotoxin induces DNA damage, cell cycle arrest, senescence and death in eukaryotic cells (Nougayrède *et al.*, 2006; Taieb *et al.*, 2016). Here we describe a method to quantify the genotoxicity of bacteria producing colibactin following a short infection of cultured mammalian cells with colibactin producing *E. coli*.

Keywords: Colibactin, Infection, *Escherichia coli*, Genotoxin, DNA damage, γ -H2AX

[Background] The genotoxin colibactin is a polyketide nonribosomal peptide hybrid compound produced by several species of *Enterobacteriaceae*. This toxin, synthesized by a machinery encoded on a 54 kb genomic locus, the *pks* island, induces DNA damages, cell cycle arrest, senescence and death in eukaryotic cells (Nougayrède *et al.*, 2006; Taieb *et al.*, 2016). The genotoxic activity of colibactin is dependent on a direct host cell-bacteria interaction and cannot be recapitulated from culture supernatant, killed bacteria, or bacterial lysates instead of live bacteria. Visualization and quantification of the colibactin genotoxic effect on eukaryotic cells can be assessed by quantification of the megalocytosis phenotype (for a protocol see Bossuet-Greif *et al.*, 2017) or quantification of the double-strand DNA breaks in the host cell nucleus by a comet assay (revealing DNA fragmentation) or phosphorylation of the H2AX histone, a marker of double-strand DNA breaks. The phosphorylation of the histone H2AX is characterized as an early and sensitive reaction to genotoxic agents (Audebert *et al.*, 2010). H2AX phosphorylation was demonstrated to be 10-100 times more sensitive than the comet assay *in vitro* as well as *in vivo* (Audebert *et al.*, 2010). The quantification of phosphorylated histone H2AX (γ -H2AX) can be processed by the In-Cell Western Assay, an immunochemical assay that uses fluorescence to detect and quantify proteins in fixed cells (Audebert *et al.*, 2010). Here we describe an adapted assay allowing the measurement of γ -H2AX in 96-well plate using In-Cell Western, following a short infection of cultured mammalian cells with colibactin-producing bacteria (Martin *et al.*, 2013; Bossuet-Greif *et al.*, 2016; Tronnet *et al.*, 2017).

Materials and Reagents

1. Black tissue culture plate 96 wells flat bottom (Greiner Bio One International, catalog number: 655090)
2. Parafilm
3. Aluminum foil
4. *pks⁺* and *pks⁻* *Escherichia coli* strains (stored in LB 20% glycerol at -80 °C)
Note: Strains typically used as positive controls in the authors' laboratory are probiotic strain Nissle 1917 or the commensal strain M1/5. Strains used as a negative control are the K-12 strain MG1655. Our lab can provide these strains.
5. HeLa cells (ATCC, catalog number: CCL-2), 20 passages maximum
6. Lennox L broth base (LB medium; Thermo Fisher Scientific, Invitrogen™, catalog number: 12780029)
7. Dulbecco's modified Eagle medium (DMEM) with 25 mM HEPES (Thermo Fisher Scientific, Gibco™, catalog number: 42430)
8. Hanks' balanced salt solution (HBSS; Sigma-Aldrich, catalog number: H8264)
9. Gentamicin solution 50 mg/ml (Sigma-Aldrich, catalog number: G1397)
10. Dulbecco's phosphate buffered saline (PBS; Sigma-Aldrich, catalog number: D8537)
11. Dulbecco's modified Eagle medium (DMEM), high glucose, GlutaMax Supplement, pyruvate (Thermo Fisher Scientific, Gibco™, catalog number: 31966021)
12. Fetal bovine serum (FBS; Thermo Fisher Scientific, Gibco™, catalog number: 10270106)
13. Non-essential amino acids solution (NEAA) 100x (Thermo Fisher Scientific, Gibco™, catalog number: 11140035)
14. Paraformaldehyde (PFA) 20% (Electron Microscopy Sciences, catalog number: 15713)
15. 10x PBS (Sigma-Aldrich, catalog number: D1408)
16. Ammonium chloride (NH₄Cl; BioUltra, Sigma-Aldrich, catalog number: 09718-250G)
17. Triton™ X-100 (Sigma-Aldrich, catalog number: X100-500ML)
18. MAXblock blocking medium (Active Motif, catalog number: 15252)
19. Phosphatase inhibitor PHOSTOP (10x, Roche Diagnostics, catalog number: 04906837001)
20. RNase (Sigma-Aldrich, catalog number: R6513)
21. Sodium chloride (NaCl; Sigma-Aldrich, catalog number: S7653)
22. Sodium azide (Sigma-Aldrich, catalog number: S8032)
23. Rabbit monoclonal anti- γ -H2AX antibody (Cell Signaling Technology, catalog number: 9718)
24. IRDye™ 800CW-conjugated goat anti-rabbit secondary antibody (2 mg/ml, Biotium (distributor Interchim), catalog number: 20078)
25. RedDot™2 (200x in DMSO, Biotium (distributor Interchim), catalog number: 40061)
26. HeLa cell culture medium (see Recipes)
27. Fixation solution (see Recipes)
28. Neutralization solution (see Recipes)

29. 10% Triton X-100 (see Recipes)
30. Permeabilization solution (see Recipes)
31. Blocking solution (see Recipes)
32. Permeabilization solution (see Recipes)
33. PST buffer (see Recipes)
34. 100x azide (see Recipes)
35. Anti- γ -H2AX solution (see Recipes)
36. Secondary antibody solution (see Recipes)

Equipment

1. 37 °C, 5% CO₂ incubator for cell cultures
2. 37 °C incubator for bacterial cultures
3. Microplate reader with 680 and 800 nm channels (here, Odyssey Infrared Imaging Scanner, Li-Cor ScienceTec, Les Ulis, France)
4. Microplate reader for absorbance measurement at 600 nm (TECAN Infinite Pro)
5. Colorimeter to measure the absorbance at 600 nm of bacterial cultures (Biochrom, model: WPA CO7500)
6. Variable Speed Rocker (VWR, catalog number: 75832-308)
7. Chemical safety hood
8. Clean bench
9. Inverted microscope (Olympus, model: CKX31)
10. Pipettes and multichannel micropipette 30-300 μ l

Software

1. GraphPad Prism 6.0

Procedure

Day 1

1. Bacteria are cultured overnight in 3 ml of LB from -80 °C glycerol stock or a colony on an LB agar plate at 37 °C with shaking (220 rpm).
2. HeLa cells are dispensed in a black 96-well tissue culture plate with transparent bottom (1.5 x 10⁴ cells/well) and grown for 24 h in 200 μ l HeLa cell culture medium (see Recipes) at 37 °C in a CO₂ incubator.

Note: Importantly, don't dispense any cells in the two bottom-left wells (see Figure 1: plate infection scheme).

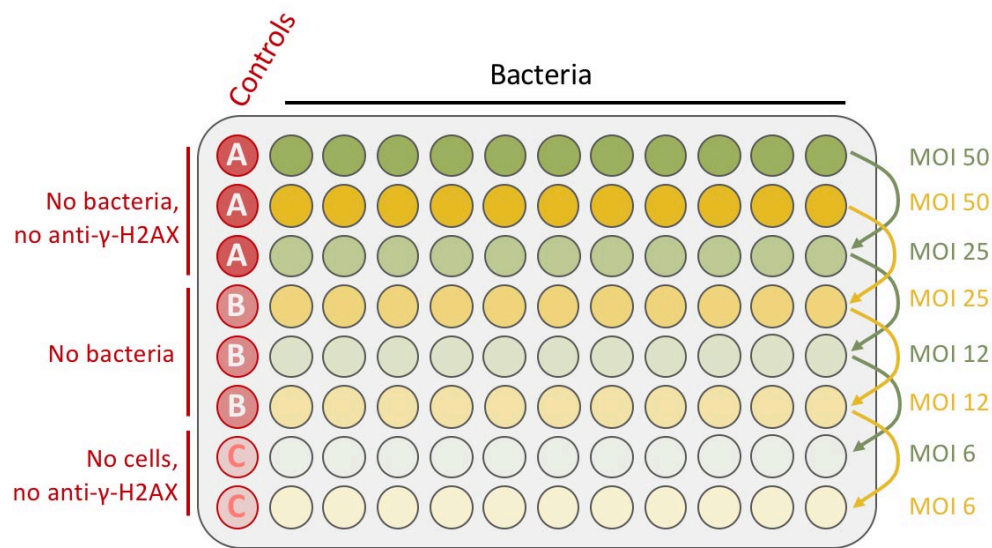


Figure 1. Schematic representation of plate infection example. Bacteria are inoculated in the two first rows in duplicate (in 200 μ l/well of DMEM-HEPES medium), at an MOI of 50 and then 2-fold serially diluted (by successively transferring 100 μ l to the rows below). The final volume is 100 μ l. The left column is kept for the controls (red wells column).

Day 2

1. Bacterial preparation

Bacteria are sub-cultured by diluting 50 μ l of the overnight culture in 5 ml of DMEM-HEPES medium (*i.e.*, 1:100 dilution) and grown at 37 °C with agitation (220 rpm) until an average $OD_{600nm} = 0.6$. The number of bacteria per ml is determined by measuring the OD_{600nm} of 1 ml of the culture.

Note: The typical value for E. coli is 1 unit of the OD_{600nm} corresponds to 5×10^8 bacteria/ml.

2. HeLa cells preparation

- Before infection, eukaryotic cells are cautiously washed once with pre-warmed (37 °C) HBSS.
- 100 μ l of DMEM-HEPES medium is added in all the wells, except the 2 first rows, corresponding to the highest number of bacteria per cell (maximum multiplicity of infection or MOI) where 200 μ l of DMEM-HEPES medium is added (Figure 1).

3. Infection assay

- After determination of the multiplicity of infection ($MOI = \text{number of bacteria per HeLa cell at the onset of the infection}$), the appropriate number of bacteria is added in the first 2 rows of the plate (containing 200 μ l DMEM-HEPES medium).
- 2-fold serial dilution is performed to create an infectious dose-effect, in a final DMEM-HEPES medium volume of 100 μ l (Figure 1).

Note: As an example, for an MOI of 50, add $50 \times 1.5 \times 10^4 = 75 \times 10^4$ bacteria for a well containing 1.5×10^4 HeLa cells. The number of dispensed cells is taken into account for the MOI calculation.

- c. The plate is incubated for 4 h in a cell culture incubator.
4. End of the infection and post-infection incubation
 - a. After infection of 4 h, the absorbance at 600 nm is measured with a microplate reader, to monitor bacterial growth. The typical values obtained are around 0.2-0.3 for MOI 50.
 - b. Then, cells are washed carefully at least 3 times with an increasing volume of pre-warm (37 °C) HBSS (100 μ l, 150 μ l and finally 200 μ l). The plate is observed under an inverted microscope to check whether most of the bacteria are removed. If required, additional washes can be performed.
 - c. Finally, cells are incubated in 200 μ l cell culture medium (see Recipes) for 3 h with 200 mg/ml gentamicin in a cell culture incubator.
5. Cells fixation
 - a. The cells are washed carefully in cold PBS (100 μ l) three times.
 - b. 100 μ l fixation solution (see Recipes) is added for 20 min under a chemical safety hood.
6. Paraformaldehyde neutralization
 - a. After the fixation step, cells are washed with PBS (100 μ l per well), for 5 min under rapid agitation (variable speed shaker, 20-25 rpm). This washing step is repeated 3 times.
 - b. Then, the paraformaldehyde is neutralized by adding 50 μ l of neutralization solution (see Recipes) for 2 min under slow agitation (3-5 rpm).
7. Cells permeabilization
 - a. The cells are washed 3 times for 5 min with PBS, 100 μ l per well, under rapid agitation (20-25 rpm) as previously (Step 6).
 - b. Then cells are permeabilized with cold permeabilization solution (see Recipes), 50 μ l per well for 5 min, under slow agitation.
8. Blocking step
 - a. Cells are washed 3 times with PST buffer (see Recipes), 100 μ l per well, under agitation (20-25 rpm).
 - b. Then, 50 μ l/well of blocking solution (see Recipes) is added, and the plate is incubated at room temperature for 60 min under agitation (3-5 rpm).
9. Anti- γ -H2AX immunostaining step
 - a. Before immunostaining, cells are washed with PST buffer 3 times, 100 μ l for 5 min under agitation (20-25 rpm).
 - b. 25 μ l per well of anti- γ -H2AX solution (see Recipes) is added and the plate is incubated for 2 h at room temperature or overnight at 4 °C (in that case, the plate has to be sealed carefully with Parafilm to avoid drying), under agitation (3-5 rpm).

Note: This solution is not added in the controls A and C (Figure 1), 25 μ l of PST buffer is added instead during the incubation time.
10. Secondary detection step
 - a. Cells are first washed 3 times with PST buffer, 100 μ l per well for 5 min under agitation (20-25 rpm).

- b. Secondary detection is carried out using an infrared fluorescent dye conjugated IRDye™ 800CW-conjugated goat anti-rabbit antibodies absorbing at 800 nm (1/1,000) in PST buffer. For DNA labeling, 1/1,000 dilution of RedDot™2 in PST is used together with the secondary antibody (see secondary antibody solution in Recipes), 25 μ l per well in every well of the plate.
- c. The plate is incubated in the dark for 1 h under agitation (3-5 rpm).

11. Plate scanning step

- a. The plate is washed 3 times in PST buffer, 100 μ l per well for 5 min under agitation (20-25 rpm) in obscurity (covered with aluminum foil).
- b. Then, the plate is dried by reversing and patting it delicately on a tissue.
- c. The DNA and the γ -H2AX were simultaneously visualized using an Odyssey Infrared Imaging Scanner with the 680 nm fluorophore (red color) and the 800 nm fluorophore (green dye), respectively (Figure 3A).

Note: Once the plate dried (Step 11b), it can be conserved several months, at 4 °C protected from the light, upside down, before visualization.

Data analysis

A quantitative analysis of the frequency of DNA double-strand breaks can be performed using the fluorescence values (in RFU) obtained from the scans (Figures 2 and 3B). Firstly, the average γ -H2AX fluorescence per cell is obtained by dividing the measured γ -H2AX fluorescence per well (at 800 nm) by the corresponding measured fluorescence for RedDot-labelled DNA per well (at 680 nm) (Figure 2). The value for γ -H2AX fluorescence per cell is divided by the average fluorescence value at 800 nm obtained for the two wells serving as vehicle control (here corresponding to the wells marked 'C' in Figures 1 and 2) to determine the percent change in phosphorylation of H2AX levels relative to the vehicle control. The obtained number is defined as a genotoxic index (or γ -H2AX fold induction) and typically ranges from 0 to 2 in completely healthy cells, to 5 to 7 in cells infected with colibactin-producing *E. coli* at MOI 25 (Figure 2 and Figure 3B).

		RED (680 nm)					Green (800 nm)		
		Controls	<i>E. coli pks+</i>	<i>E. coli pks-</i>			Controls	<i>E. coli pks+</i>	<i>E. coli pks-</i>
Raw values	A	18,58	12,73	19,07	Raw values	A	1,18	120,04	23,19
	B	23,96	19,01	19,49		B	0,86	111,79	28,45
	C	16,6	21,31	21,34		C	1,16	100,63	23,87
	D	16,51	26,46	21,71		D	31,98	107,43	17,39
	E	16,82	21,18	19,89		E	30,29	83,95	26
	F	17,01	19,31	19,58		F	30,18	49,43	27,75
	G	2,67	19,41	20,74		G	0,44	31,75	24,11
	H	3,5	20,04	17,34		H	0,31	26,74	23,46
Background red :		3,085			Background green :		1,067		
Background subtracted		Controls	<i>E. coli pks+</i>	<i>E. coli pks-</i>	Background subtracted		Controls	<i>E. coli pks+</i>	<i>E. coli pks-</i>
	A	15,495	9,645	15,985		A	0,113	118,973	22,123
	B	20,875	15,925	16,405		B	-0,207	110,723	27,383
	C	13,515	18,225	18,255		C	0,093	99,563	22,803
	D	13,425	23,375	18,625		D	30,913	106,363	16,323
	E	13,735	18,095	16,805		E	29,223	82,883	24,933
	F	13,925	16,225	16,495		F	29,113	48,363	26,683
	G	-0,415	16,325	17,655		G	-0,627	30,683	23,043
	H	0,415	16,955	14,255		H	-0,757	25,673	22,393
Average duplicates		Controls	<i>E. coli pks+</i>	<i>E. coli pks-</i>	Average duplicates		Controls	<i>E. coli pks+</i>	<i>E. coli pks-</i>
	A-C	16,628	12,785	16,195		A-C	0,00	114,848	24,753
	D-F	13,695	20,8	18,44		D-F	29,75	102,963	19,563
			17,16	16,65				65,623	25,808
			16,64	15,955				28,178	22,718
		<i>E. coli pks+</i>					<i>E. coli pks-</i>		
		Ratio : $\frac{[(A-B)]}{[(A-B)]}$					Ratio : $\frac{[(A-B)]}{[(A-B)]}$		
		Genotoxic index: Ratio/Ctrl					Genotoxic index: Ratio/Ctrl		
Ctrl (without bacteria) $\frac{[(D-F)]}{[(A-C)]}$:		1,789114964							
	MOI 25	A-B	8,983053057	5,020947919	MOI 25	A-B	1,528455285	0,85430803	
	MOI 12	C-D	4,950160256	2,766820665	MOI 12	C-D	1,060918294	0,59298498	
	MOI 6	E-F	3,824203574	2,137483421	MOI 6	E-F	1,55005005	0,86637811	
	MOI 3	G-H	1,693409455	0,946506786	MOI 3	G-H	1,423900554	0,79586867	

Figure 2. Data analysis of γ -H2AX quantification. Representative image of how we process data from the raw numbers to the calculation of the γ -H2AX fold induction (genotoxic index).

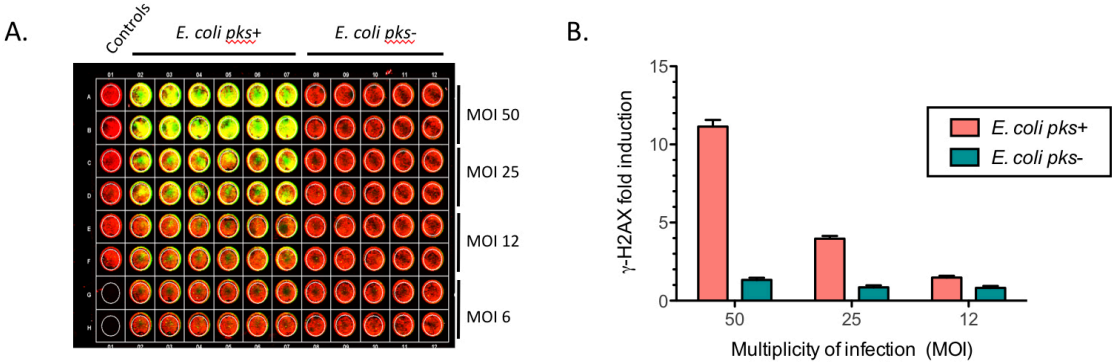


Figure 3. Quantification of phosphorylated H2AX (γ -H2AX) from infected HeLa cells with *pks+* *E. coli* or with *pks-* *E. coli* at various MOI. A. Example of an In-Cell Western plate image with merged detection of total DNA (red, 680 nm) and γ -H2AX (green, 800 nm). A high level of γ -H2AX can be observed at high MOI for the cells infected with *E. coli pks+*. B. The γ -H2AX fold induction (γ -H2AX fluorescence normalized to the amount of cells per well), calculated relative to the control (non-infected cells), reveals a genotoxic dose-response depending on the MOI in cells infected by colibactin-producing bacteria (*E. coli pks+*) compared to cells infected by non colibactin-producing bacteria (*E. coli pks-*). Data represented in the graph were obtained from two biological replicates and two independent experiments. Data were plotted using GraphPad Prism 6.0. The mean with standard deviation (sd) is shown.

Recipes

1. HeLa cells culture medium
450 ml DMEM high glucose with pyruvate and GlutaMax
50 ml fetal bovine serum
5 ml NEAA
2. Fixation solution (1x PBS, 4% PFA)
5 ml PFA stock solution
5 ml 20% 10x PBS
40 ml distilled water
Aliquot and store at -20 °C
Defreeze one aliquot the day of the experiment
3. Neutralization solution (20 mM NH₄Cl in PBS)
 - a. Prepare the stock solution (1 M NH₄Cl) in distilled water and store at 4 °C
 - b. Dissolve the stock solution in PBS to reach the appropriate concentration
 - c. Keep on ice until use
4. 10% Triton X-100
500 μ l Triton X-100 in 50 ml PBS
This solution can be stored at 4 °C for 2-3 weeks
5. Permeabilization solution (1x PBS, 0.2% Triton X-100)
200 μ l 10% Triton X-100
10 ml PBS
Keep on ice until use
6. Blocking solution (MAXblock blocking medium, 1x PHOSTOP inhibitor, RNase [100 μ g/ml])
4.5 ml MAXblock blocking medium
500 μ l 10x PHOSTOP inhibitor
50 μ l RNase stock solution (10 mg/ml in PBS, 10 mM HEPES pH 7.5, 15 μ M NaCl, aliquot stored at -20 °C)
Keep on ice until use
7. PST buffer (PBS, 2% fetal calf serum, 0.2% Triton X-100)
48 ml PBS
1 ml 2% fetal calf serum
1 ml 0.2% Triton X-100 in PBS
8. 100x azide
Dissolve sodium azide in distilled water at a final concentration of 50 mg/ml (5% w/w), store at 4 °C
9. Anti- γ -H2AX solution (rabbit monoclonal anti- γ -H2AX [1/200] in PST buffer, 500 μ g/ml azide)
1.5 ml PST buffer
7.5 μ l anti- γ -H2AX

15 μ l 100x azide

Note: This solution can be reused 2-3 times, and stored at 5 °C for several weeks.

10. Secondary antibody solution (IRDye™ 800CW Goat Anti-Rabbit [1/1,000], RedDot™2 [1/1,000] in PST buffer)

2.5 ml PST

2.5 μ l IRDye™ 800CW Goat Anti-Rabbit

2.5 μ l RedDot™2

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References

1. Audebert, M. Riu, A., Jacques, C., Hillenweck, A., Jamin, E. L., Zalko, D., Cravedi, J. P. (2010). [Use of the gammaH2AX assay for assessing the genotoxicity of polycyclic aromatic hydrocarbons in human cell lines](#). *Toxicol Lett* 199: 182-192.
2. Bossuet-Greif, N., Belloy, M., Boury, M., Oswald, E. and Nougayrede, J. (2017). [Protocol for HeLa cells infection with *Escherichia coli* strains producing colibactin and quantification of the induced DNA-damage](#). *Bio-protocol* 7(16): e2520.
3. Bossuet-Greif, N., Dubois, D., Petit, C., Tronnet, S., Martin, P., Bonnet, R., Oswald, E. and Nougayrede, J. P. (2016). [Escherichia coli C1bS is a colibactin resistance protein](#). *Mol Microbiol* 99(5): 897-908.
4. Martin, P., Marcq, I., Magistro, G., Penary, M., Garcie, C., Payros, D., Boury, M., Olier, M., Nougayrede, J. P., Audebert, M., Chalut, C., Schubert, S. and Oswald, E. (2013). [Interplay between siderophores and colibactin genotoxin biosynthetic pathways in *Escherichia coli*](#). *PLoS Pathog* 9(7): e1003437.
5. Nougayrède, J. P., Homburg, S., Taieb, F., Boury, M., Brzuszkiewicz, E., Gottschalk, G., Buchrieser, C., Hacker, J., Dobrindt, U. and Oswald, E. (2006). [Escherichia coli induces DNA double-strand breaks in eukaryotic cells](#). *Science* 313(5788): 848-851.
6. Taieb, F., Petit, C., Nougayrede, J. P. and Oswald, E. (2016). [The enterobacterial genotoxins: cytolethal distending toxin and colibactin](#). *EcoSal Plus* 7(1).
7. Tronnet, S., Garcie, C., Brachmann, A. O., Piel, J., Oswald, E. and Martin, P. (2017). [High iron supply inhibits the synthesis of the genotoxin colibactin by pathogenic *Escherichia coli* through a non-canonical Fur/RyhB-mediated pathway](#). *Pathog Dis* 75(5).