



Report



AUSTRIAN
MARSHALL PLAN FOUNDATION

Effects of *Lactobacillus johnsonii* on genotoxicity after radiation
treatment in mice

(Within the project title: Influence of gut microbiota on B-lymphocytes genotoxicity in
Ataxia telangiectasia patients)

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Acknowledgment

First, I would like to thank Dr. Robert Schiestl for the great opportunity to work in his research group in the Environmental Health Science department at the UCLA Fielding School of Public Health and Dr. Irene Maier for her wonderful guidance and advice during my stay. It was a unique chance for me to take part in many exciting discussions and to gain a comprehensive insight into his research.

Special thanks to the Austrian Marshall Plan Foundation for the generous financial support, without this stay would not have been possible. Moreover, I highly appreciate the financial support from the University of Vienna.

Further I would also like to thank the whole group of Robert Schiestl for including me so kindly, giving me support and positive input in all issues and making this time unforgettable for me. Special thanks to Chayo Minutti for dealing with all the paper work, Barbara Housel for helping me with the university application process, Liuba Parfenova and Zorica Scuric. It was very impressive to work in such an outstanding group.

And finally, I want to thank Prof. Dr. Karl-Heinz Wagner for his professional help and support during my yearlong application process.

Table of contents

Acknowledgment.....	2
Table of contents	3
List of Figures	5
List of Tables.....	6
Abstract.....	7
1 Literature.....	8
1.1 The intestine	8
1.2 Gut microbiota	8
1.2.1 Composition and development of the intestinal microbiota	9
1.2.2 Gut homeostasis - microbiota in health and disease.....	10
1.3 Immune system, intestinal inflammation and genotoxicity	12
1.3.1 The immune system	12
1.3.2 Inflammation and genotoxicity	13
1.3.3 Resulting questions.....	19
1.4 Selection of probiotic bacterial strains	19
1.4.1 <i>Lactobacillus johnsonii</i>	20
1.5 Mouse models	22
2 Methods and Material	24
2.1 Animals and husbandry conditions	24
2.2 Cultivation of <i>Lactobacillus johnsonii</i> and inoculation of mice.....	24
2.2.1 Growing <i>Lactobacillus johnsonii</i>	25
2.2.2 Inoculating the mice	25
2.2.3 Reagents for bacterial growing and inoculation.....	25
2.2.4 Material for bacterial growing and inoculation	25

2.3	Isolation of Intraepithelial Lymphocytes from mice	26
2.3.1	Protocol	26
2.3.2	Material for Isolation of Intraepithelial Lymphocytes from mice	28
2.3.3	Reagents for Isolation of Intraepithelial Lymphocytes from mice ..	28
2.4	γ H2AX foci determination on Blood Lymphocytes	29
2.4.1	Background and principles of γ H2AX assay	29
2.4.2	Protocol for γ H2AX assay	32
2.4.3	Material for γ H2AX assay	34
2.4.4	Reagents for γ H2AX assay	35
2.5	Micronucleus determination in Blood Erythrocytes	35
2.5.1	Background and principles of micronucleus formation	35
2.5.2	Protocol for micronucleus determination	39
2.5.3	Material for micronucleus determination.....	39
2.5.4	Reagents for micronucleus determination	40
2.6	Single cell gel electrophoresis (comet assay) for lymphocytes from whole blood	40
2.6.1	Background and general principles of the alkaline comet assay ...	40
2.6.2	Methodology – step by step	43
2.6.3	Solution preparation for comet assay	45
2.6.4	Protocol for comet assay	46
2.6.5	Material for comet assay	48
2.6.6	Reagents for comet assay.....	48
3	References	49

List of Figures

Figure 1 concentration and distribution of commensal intestinal microbiota	10
Figure 2 Latitudinal distribution from the epithelial surface to intestinal lumen .	10
Figure 3 balance of a healthy gut homesostasis	11
Figure 4 Diet, microbial composition and regulation of the immune system.. ...	14
Figure 5 Overview of the cellular signaling pathways	17
Figure 6 Potential inflammatory mechanisms	18
Figure 7 harvesting intraepithelial lymphocytes	26
Figure 8 γ H2AX phosphorylation	29
Figure 9 Organisation of DNA and γ H2AX foci	30
Figure 10 Principle of γ H2AX assay and foci pictures.....	31
Figure 11 Erythropoiesis	37
Figure 12 micronuclei with Wright's Giemsa staining.....	39
Figure 13 General principle of comet assay.....	41
Figure 14 Fluorescent pictures of comets under the microscope.....	41

List of Tables

Table 1 Criteria of properties for a good probiotic strain	20
Table 2 Reagents for bacterial growing and inoculation	25
Table 3 Material for bacterial growing and inoculation	26
Table 4 Material for Isolation of Intraepithelial Lymphocytes from mice	28
Table 5 Reagents for Isolation of Intraepithelial Lymphocytes from mice	28
Table 6 Material for γ H2AX assay	34
Table 7 Reagents for γ H2AX assay	35
Table 8 Material for micronucleus determination	39
Table 9 Reagents for micronucleus determination	40
Table 10 Material for comet assay	48
Table 11 Reagents for comet assay	48

Abstract

Within the last decade there has been an increasing interest to identify microbial inhabitants in the gastrointestinal tract of humans and to understand their beneficial and detrimental role in health and disease. The gut microbiota – a symbiosis between the host and the microbes - plays an important role in human metabolism and on the other hand the bacteria benefits from the nutrient-rich niche in the intestine. Perhaps even more crucial is that the gut microbiota interacts with the human immune system, by stimulating signaling pathways to promote the maturation of immune cells and to activate other immune functions. However, if the gut homeostasis is out of balance, chronic inflammation in the gut can lead to increased DNA damage which is highly associated with diseases like obesity, type 1 diabetes, inflammatory bowel disease including Crohn's disease and colitis ulcerosa, asthma and colon cancer.

Previous results show that several *Lactobacillus johnsonii* strains have advantageous health effects when used as probiotic strain in human and animal administration.

In this study we investigated two groups of mice with a different composition of the intestinal microbiota (conventional vs. restricted) and husbandry conditions (pathogen-free vs. sterile) and collected data of inflammatory response due to *L.johnsonii* administration on genotoxicity.

1 Literature

1.1 The intestine

The human gut with its surface of approximately 200 to 300m² represents a major area of exogenous environmental impact to the human body. The gut-associated lymphoid tissue (GALT) makes the gut the biggest and most important immunological organ. (Collins et al.; 1998) The gastrointestinal system is not only here for digesting food, absorption of nutrients and excreting indigestible compounds but likewise has a huge impact on the immune system due to the immense microbiological colonization. It is from greatest importance that our body has the ability to distinguish between beneficial and pathogenic compounds to achieve a healthy lifestyle. (Bischoff; 2011)

1.2 Gut microbiota

Within the last decade there has been an increasing interest to identify microbial inhabitants such as bacteria, archaea, viruses and unicellular eukaryotes in the gastrointestinal tract of humans and to understand their beneficial and detrimental role in health and disease. The commensal gut microbiota – a symbiosis between the host and the microbes - plays an important role in human metabolism and on the other hand the bacteria benefits from the warm, humid and nutrient-rich niche in the intestine (Chung et al.; 2010). As mentioned above, the gut microbiota interacts with the human immune system, by stimulating signaling pathways to promote the maturation of immune cells and to activate other immune functions (Sartor; 2008; Clemente et al.; 2012). Some years ago, the Human Microbiome Project was initiated to analyze and sequence the huge diversity of microbes in our intestine. It is estimated that the total human microbiota on skin, gastrointestinal tract and respiratory tracts, contains about 10¹⁴ bacterial cell which is 10 times more than the number of cells present in the human body. The gut is by far the most colonized and at which the colon contains approximately 70% of all microbes (Ley et al.; 2006). This gives an idea about how powerful the interaction between the host and the microbiota can be.

1.2.1 Composition and development of the intestinal microbiota

Immediately during birthing process the baby gets into contact with the mothers microflora and for the first time it is exposed to a complex microbial population. Within the first year of life the composition of the child's microbiota develop and hence the immune system. However, after its first year the essential composition begins to stabilize. A comparison between parents and their children shows similarity and therefore the parent's gut microbiota can be seen as a major factor in shaping the intestinal community of their offspring. (Mandar et al.; 1996)

Besides the maternal influence diet, physiological aspects, environmental exposure, pathogens, competition within the resident bacteria and ability of adaptation as well as antimicrobial therapies and host genetics are the most important factors developing the commensal flora over the years. Every human has his/her own individual composition and is dependent on the factors mentioned before. (Ley et al.; 2006; Sekirov et al.; 2010)

16S ribosomal RNA gene sequence based molecular analysis of human fecal and mucosal samples detected around 36.000 different species. All microorganisms together contain at least 100 times as many genes as the human genome. (Sartor et al.; 2012) More than 99% of the intestinal microbiota originates from four bacterial divisions: *Firmicutes*, *Bacteroidetes*, *Proteobacteria* and *Actinobacteria* whereas *Firmicutes* and *Bacteroidetes* together represent already 90%. (Qin et al.; 2010) Both, complexity and concentration increase from stomach (10^2) to colon (10^{12}) (Fig. 1). In addition to the longitudinal heterogeneity there is also a big latitudinal variation in the microbiota composition (Fig. 2). The latitudinal compartment can be roughly divided into three habitats the epithelial surface, mucus layer and intestinal lumen and in each of those can be found different strains. (Swidsinski et al.; 2005)

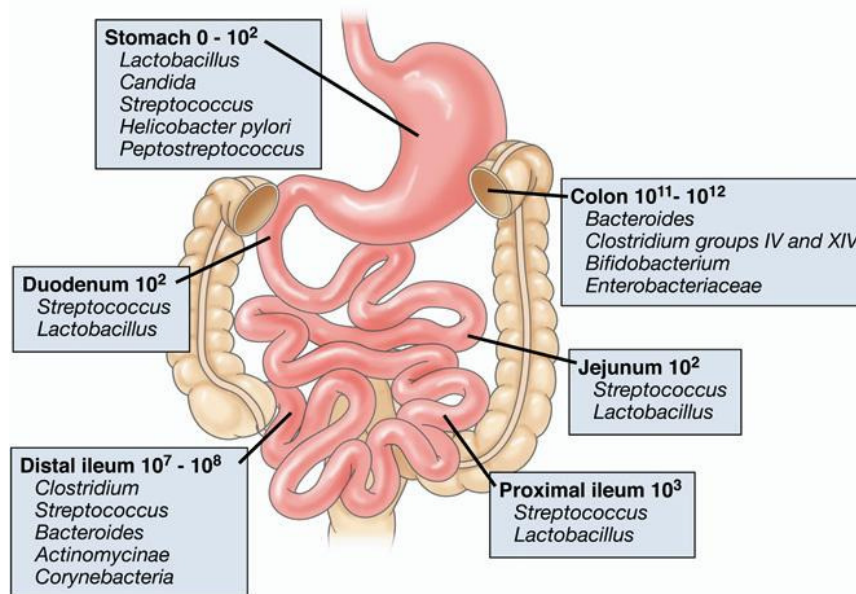


Figure 1 concentration and distribution of commensal intestinal microbiota

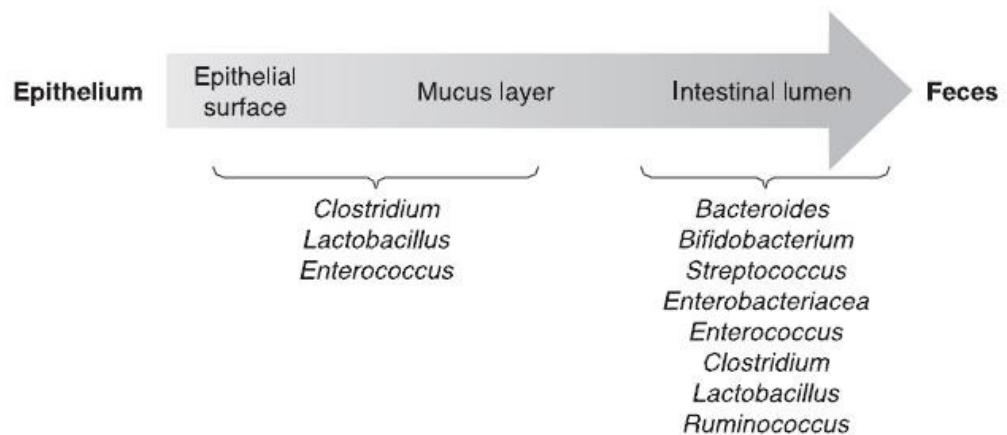


Figure 2 Latitudinal distribution from the epithelial surface to intestinal lumen

1.2.2 Gut homeostasis - microbiota in health and disease

The history of co-evolution between mammals and microbiota is long. Already in ancient times the gut wellbeing got attention, where 400 B.C. Hippocrates said “death sits in the bowels” (Hawrelak et al.; 2004). A healthy host has the ability to control resident commensal bacteria without an adverse immune response. The gut homeostasis can be seen as a two-way dialogue (Fig. 3):

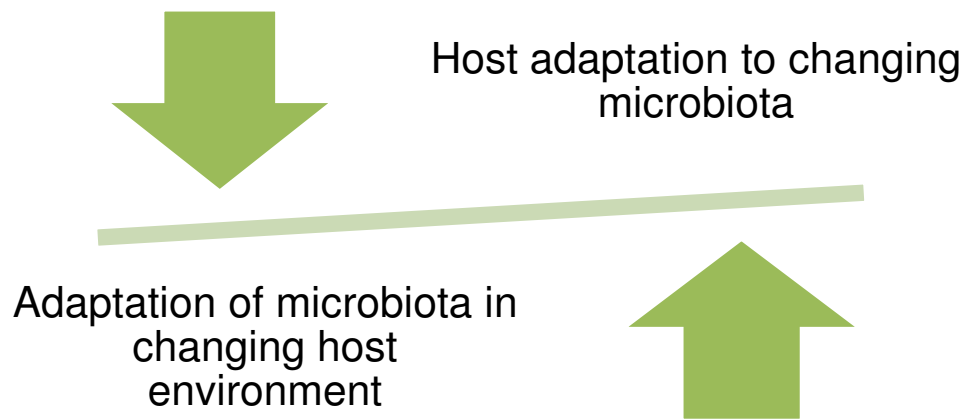


Figure 3 balance of a healthy gut homeostasis

After the common understanding that gut health and disease is not only dependant on pathogenic bacteria but rather on intestinal inhabitants an increase in research on gut microbiota was noticed. Different models both in mice and human were established to extract particular bacteria and discover their functions. Later in this chapter this will be discussed in more detail.

As mentioned before infants develop their general microbial composition within their first year. At the same time symbiotic bacteria optimize nutrient absorption, promote growth and healing, induce angiogenesis, develop the immune system and soothe inflammation (Greer et al.; 2011). Further, nonpathogenic bacteria accomplish many beneficial functions such as synthesis of vitamins, digesting fiber, antagonize pathogenic bacteria and the regulation of inflammatory response (Maslowski et al.; 2011).

Because of a changing lifestyle called Westernization this homeostasis system gets out of balance. Basically this is due to a changing diet with reduced intake in complex carbohydrates, an increase in animal products, underused breastfeeding and antibiotic medication. This leads to increased immune response caused by altered gut microbial composition which is highly associated with obesity, type 1 diabetes, inflammatory bowel disease including Crohn's disease and colitis ulcerosa, asthma and colon cancer. (Chung et al.; 2010; Greer et al.; 2011)

1.3 Immune system, intestinal inflammation and genotoxicity

The immune response consists of an enormous complex signaling pathway system which will be partly described in this chapter. A short overview should give a basic knowledge of the regulation of the immune system which is essential to understand this study and the used methods. Further, the communication of the intestinal microbiota and host will be shown and detrimental aspects on health and disease will be discussed. This essential co-existence can be divided into 4 main categories of interaction:

- Signaling Between the Microbiota and the Host
- Signaling Between the Microbiota and Pathogens
- Signaling Between Members of the Microbiota
- Signaling Between the Host and Pathogens

1.3.1 The immune system

The immune system consists of the innate and the adaptive immune system and at which the innate immune system provides an immediate but nonspecific response and the adaptive immune system also called specific immune defense is specific to antigens from bacteria, viruses, fungi, parasites and their presenting or producing macromolecules. Latest leads to the production of **T- and B- lymphocytes**, both develop from hematopoietic stem cells in the bone marrow. B-lymphocytes belong to the humoral immune system whereas T-lymphocytes are involved in the cell-mediated immune system. The reason why our immune system develops and makes it more effective is because of the memory effect. The memory effect describes the process of transforming antigen-activated T- lymphocytes into long-life memory cells with individual receptors which stay in lymphoid tissue, in mucosal barriers and in the circulation after an infection. (Silbernagl et al.; 2003)

How do B- and T-lymphocytes work? So-called antigen presenting cells (APC) such as macrophages, B-cells and dendritic cells present antigens on their cell

surface. These will be recognized by either CD8⁺ or CD4⁺ “native” T-lymphocytes and interleukin (IL) 2 is released which is the signal for T- cell proliferation. If the CD8⁺ receptor is activated, killer T- cells (cytotoxic killer cells) will be produced. When killer cells find their corresponding infected cells they release apoptotic signals. From higher importance in this work are CD4⁺ T-cells which generate T_{H1} or T_{H2} helper cells. They do not kill effected cells or pathogens directly but they direct other cells to do so. The most important cytokine produced by T_{H1} helper cells is interferon- γ (IFN- γ) which is an activator of macrophages and causes inflammation. On the other hand T_{H2} helper cells are necessary to activate B- lymphocytes. B- cells present antigens from e.g. bacteria because of a previous uptake and intracellular processing step. The antigen presenting B- cells can be activated by CD4 T cell receptor from T_{H2} helper cells by segregation of IL-4 (and IL-6). After a successful cooperation B- lymphocytes start to divide and the offspring produce an enormous amount of antibodies circulating in the blood. There they will neutralize and eliminate antigens. (Silbernagl et al.; 2003; Abbas et al.; 2012)

These two types of lymphocytes (B- and T-) play an important role in the collection of immunologic data. However, these are not all of the immune regulatory cells also yet to include are granulocytes, dendritic cells and macrophages. Moreover, there are lots of subtypes of each mentioned before.

1.3.2 Inflammation and genotoxicity

What are the underlying causes of increased inflammation or reduced inflammatory response?

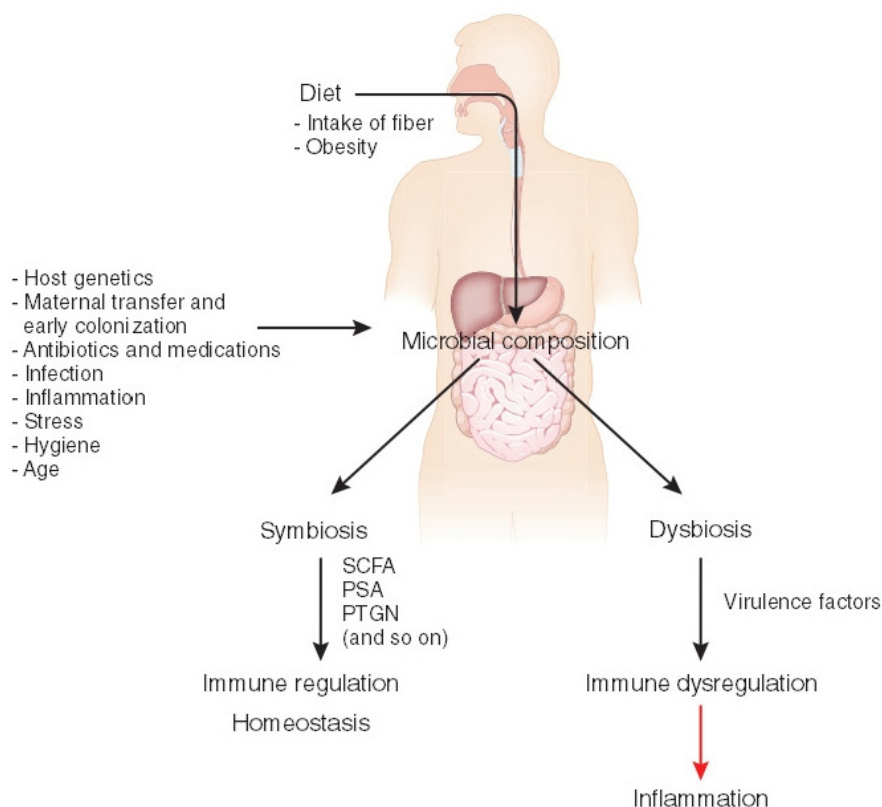


Figure 4 Diet, microbial composition and regulation of the immune system. Diet and other environmental and host factors have a major effect on intestinal gut microbiota.

The main focus will be on microbiota and also host alterations and their interaction. A minor focus will be on environmental aspects as this goes beyond to the interests of this thesis.

In healthy hosts, intestinal commensal flora activates several groups of homeostatic response either by epithelial cells, T- and B- lymphocytes, macrophages or dendritic cells to achieve a well-balanced coexistence.

Major key players in maintaining gut homeostasis which can be explained as the tolerance of antigens derived from the commensal flora or diet are **regulatory T- cells** (T_{reg}). People with a high intake of fiber show increased colonization of *Clostridium* ssp. Cluster IV and XIVa and *Bacteroides fragilis* which help to maintain gut homeostasis (Maslowski et al.; 2011). The fermentation of dietary fiber and complex O-linked mucin glycans in the mucus layer of the intestine produces large amounts of **short chain fatty acids** (SCFA). On the one hand SCFA like acetate and lactate are toxic to some

pathogens and on the other hand SCFA such as propionate and butyrate are a main nutrient source for intraepithelial cells (Maynard et al.; 2012). *B. fragilis* produce polysaccharide A from glycans which mediates the conversion of **CD4+ T- cells** into Forkhead box P3 regulatory T cells (Foxp3 T_{reg}) that produce the anti-inflammatory cytokine IL-10 (Round et al.; 2010). Therefore T_{reg} cells expressing the transcription factor Foxp3 are important to limit intestinal inflammation because IL-10 blocks the activation of **T_H17/T_H1 helper cells** (Fig 6D). A concentration range from 50 to 100mM of SCFA can be seen as sufficient (Smith et al.; 2013). Several studies on IL-10^{-/-} mice have shown depletion leads to excessive colonic inflammation and inflammatory bowel disease (IBL) (Kuhn et al.; 1993).

T_H17 cells can be seen controversial. On the one hand they are stimulated from the commensal microbiota by activating CD4⁺ T-cells via transforming growth factor β (TGF- β) and IL-6 to contribute significantly to granulopoiesis regulation, neutrophil recruitment and antimicrobial peptide (REGIII γ) induction. However, on the other hand T_H17 cell play a major role in inducing autoimmune diseases such as IBD. (Chung et al.; 2010)

The **gut epithelial barrier** is central to intestinal defenses. It is not just a simple passive barrier for microbiota and microbe-associated molecular patterns (MAMPs) such as lipopolysaccharide, peptidoglycan and flagellin but rather an active sensor for these. The epithelium develops its cells from stem cells located near the base of the intestinal crypts. Epithelial cells can directly interact with the gut lumen by releasing protective substances like mucin secretes and antimicrobial peptides or indirect basolateral by cytokines or chemokines, respectively. Intestinal epithelial cells (IECs) are supported by additional immune cells and structures including Payer's patches of the distal ileum, isolated lymphoid follicles (IFLs) and mesenteric lymph nodes – the so-called GALT. (Maynard et al.; 2012)

An important role of recognizing bacteria and MAMPs are receptors on the surface of the epithelial cells which activate the early innate immune system. (Maynard et al.; 2012) The most recent ones are from the **Toll-like receptor**

(TLR) family with different ligands for different patterns of microbial components. Ligations of these receptors stimulate the **MyD88- dependant TLR** signaling pathway (Fig. 5). MyD88 plays a major role in signal transduction and it has been shown in several studies that a loss of function leads to altered microbiota and followed by diseases. The stimulation of MyD88 induces the activation of the transcription factor **NF-kB** and AP-1 via several transduction steps (Takeda et al.; 2004). NF-kB is responsible for the transcription of both pro- and anti- inflammatory cytokines such as tumor necrose factor (TNF- α), IL-1, IL-6 and others (Sartor; 2008). However, altered NF-kB activity is linked to cancer development and progression through its ability to induce the production of adhesion molecules, reactive oxygen and nitrogen species and cyclooxygenase 2 (COX-2) (Fig.5). NF-kB can be seen as a link between inflammation and cancer (Karin; 2006). COX-2 produces prostaglandins, which are also key mediators in inflammation (Kipanyula et al.; 2013). Besides the MyD88- dependant signaling, there is a **MyD88- independent pathway** (TRIF) which releases interferon β (IFN- β). It is discussed if the MyD88- independent pathway can induce NF-kB activation. (Takeda et al.; 2004)

Though activation of the TRL pathway, repair mechanism of damaged IECs are activated and promote proliferation. Further, bacterial signals are required for the induction of antimicrobial proteins (Hooper et al.; 2012).

Next to the TRL pathway, there are series of **nucleotide-binding oligomerization domain (NOD) receptor** families. Especially NOD2(CARD15) should be mentioned at this position. This gene encodes an NOD- like receptor that is sensitive to microbiota and releases antimicrobial peptides by Paneth cells like regenerating islet- derived protein 3 γ (REGIII γ). Many of these special IECs are located in the crypts of the small intestine (Maynard et al.; 2012). A reduction of NOD2 expression leads to altered composition of bacteria for the benefit of pathogens and it was the first susceptibility gene linked to Crohn's disease. (Hugot et al.; 2001; Ogura et al.; 2001)

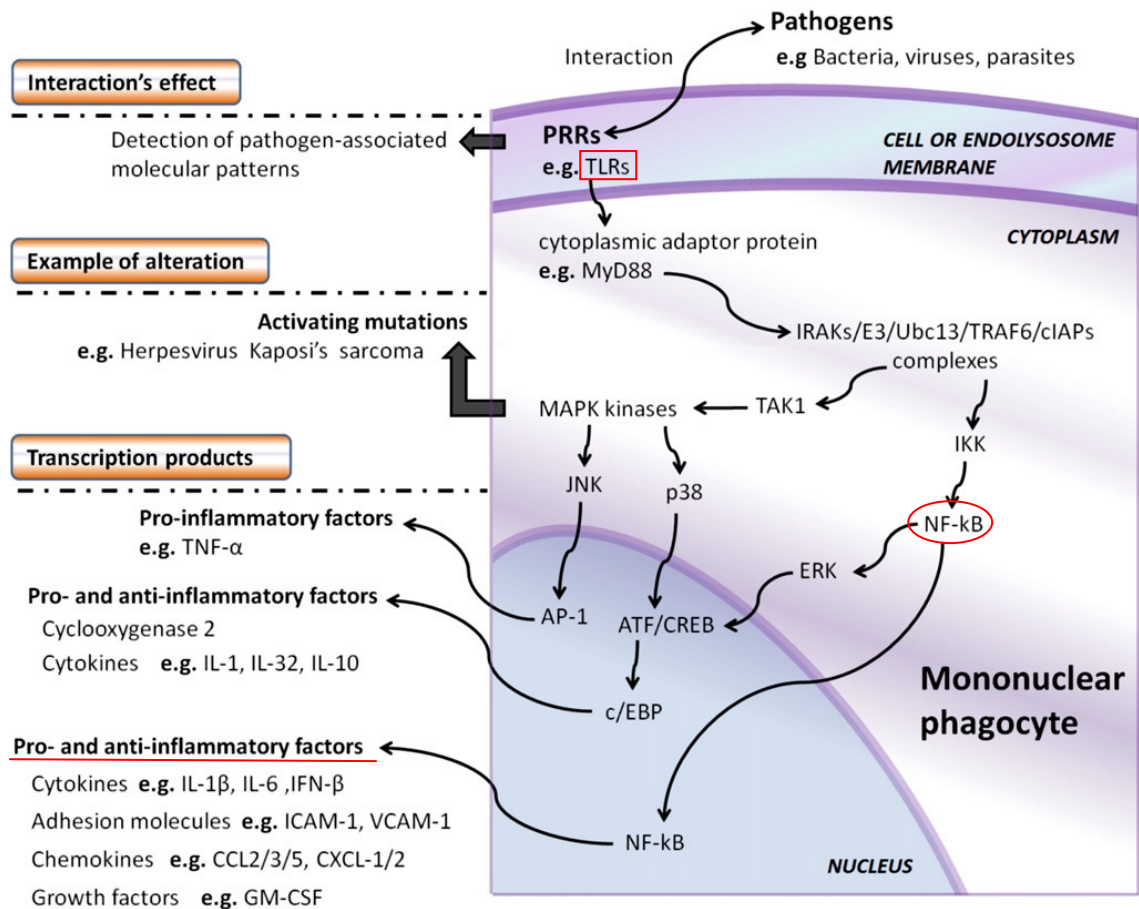


Figure 5 Overview of the cellular signaling pathways by activation of pattern recognition (PRR), like toll – like receptors (TLRs) on the epithelial surface. These pathways induce the production of either pro- or pro- and anti- inflammatory factors.

Macpherson et al. have shown that immunoglobulinA **IgA** has a protective function against pathogens and mucosal penetration by commensal microbiota. It is produced with the help of dendritic cells which absorb the bacteria from the inner mucus layer. After this procedure, activated dendritic cells interact with B-lymphocytes in the Peyer's patches, inducing IgA+ B- cells to produce IgA. This is transcytosed across the epithelium and released into the intestinal lumen (Macpherson et al.; 2004). IgAs coat antigens and commensal bacteria to inhibit their binding to the host epithelium and penetration into the lamina propria. Hence, IgA have a crucial task to achieve gut homeostasis. (Kamada et al.; 2013)

Another key player in inflammation response is the systemically circulating pro-inflammatory cytokine **tumor necrosis factor α** (TNF- α) (Fig.5). It is highly associated with persistent chronic inflammation and therefore promotes the

development of cancer. Responsible for the transcription are two receptors: tumor necrosis factor receptor 1 and 2 (TNFR1/TNFR2) (Westbrook et al.; 2012). Intestinal microbiota can activate mitogen-activated protein kinases (MAPKs) via epithelial stimulation. Further, MAPKs activate the transcription factor AP-1 and synthesize TNF- α . A proposed mechanism of TNF- α is an up-regulation of reactive oxygen and nitrogen species (RONS) which is associated with DNA damage (Chen et al.; 2008).

The interaction of the host immune system and the microbiota is remarkably complex and gives lots of possibilities for dysfunctions. The most relevant are summarized above where the following figure gives a comprehensive overview:

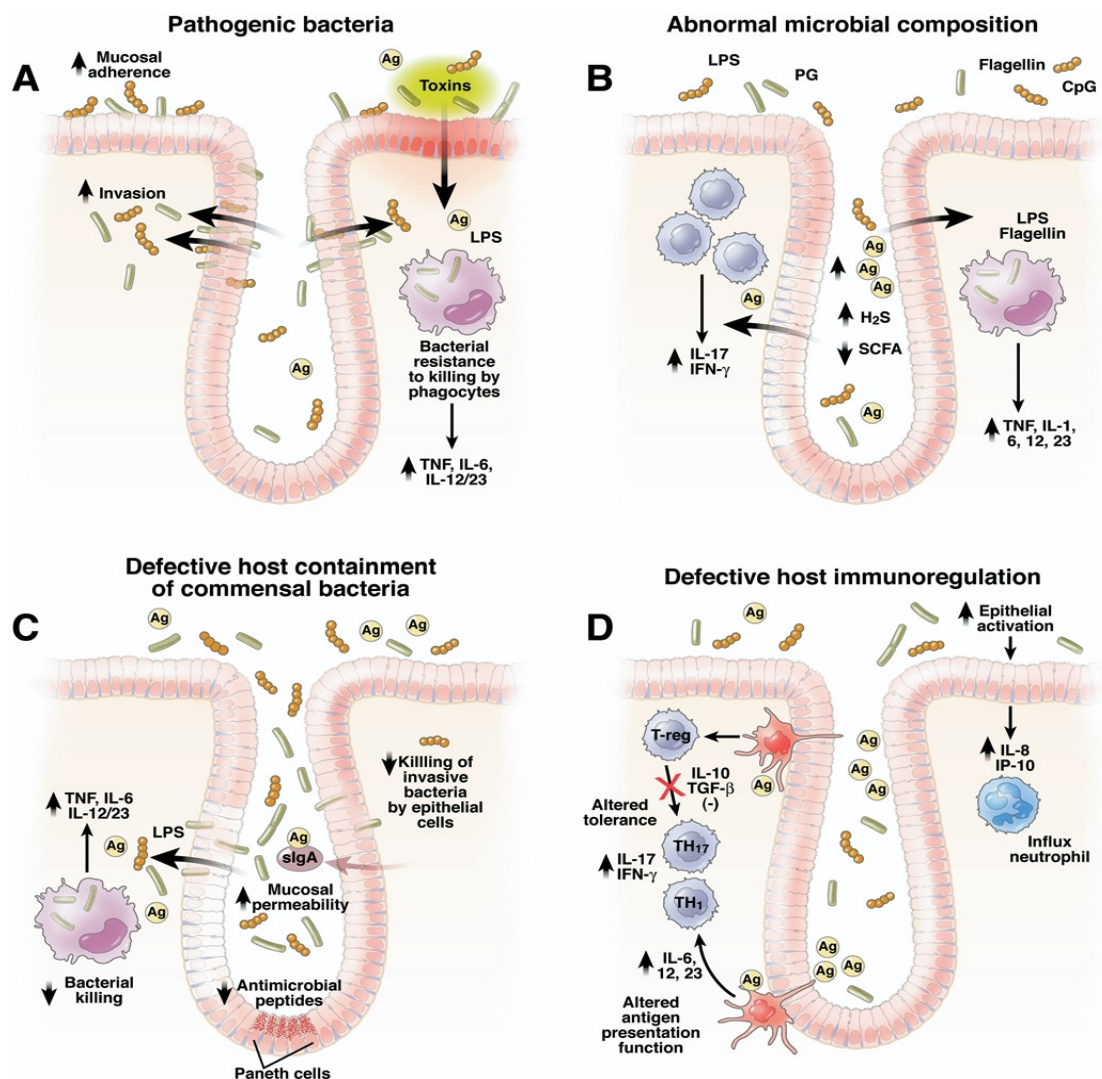


Figure 6 Potential inflammatory mechanisms by which intestinal bacteria and microbial pattern induce chronic immune-mediated intestinal and systemic injury.

1.3.3 Resulting questions

- How can we achieve a reduction of inflammation?
- Which mouse model fits best?
- How to look for relevant single strain bacteria with positive effects on gut homeostasis?
- Which methods are most suitable in order to detect alterations in inflammation and genotoxicity?

1.4 Selection of probiotic bacterial strains

According to the FAO of the UN and the WHO, probiotics are “live microorganisms” which, when administered in adequate amounts, confer a health benefit on the host.

Many scientists initiated studies to find single strains of bacteria which may have effect human health, beneficially. To detect whether there are positive or negative effects associated with a single bacterial strain it first must be isolated in pure culture, must be cultivated and has to be brought back to the intestine as viable strain (Yamano et al.; 2006). However, the majority of microbes colonizing the gut have not yet been successfully cultivated by current techniques.

Collins J.K. et al. published a list of criteria (Table 1) how to choose a good probiotic strain, because lots of papers have been published and numerous probiotic products are found on the market and therefore skepticism has risen regarding their proposed beneficial effects. Further it was not proved if these selected microorganisms are viable and have the ability survive the gastrointestinal tract. (Collins et al.; 1998)

1	Human origin
2	Possession of GRAS status (generally regarded as safety)
3	Possession of a desirable antibiogram profiles e.g. metronidazole resistance with desirable sensitivities
4	Production of antibacterial factors antagonistic for potentially pathogenic microorganisms, particularly invasive Gram negative pathogens

5	Desirable metabolic activity
6	Technological suitability
7	Non-pathogenic even in immunocompromised hosts
8	Non-inflammatory-promoting microorganisms
9	Survival in association with the adult mucosal immune system
10	Immunostimulatory for the mucosal immune system with appropriate cytokine stimulation
11	Anti-mutagenic and anti-carcinogenic properties (protection against genotoxic agents)
12	Potential vehicle for the delivery of recombinant proteins and peptides in a site specific fashion to the human gastrointestinal tract

Table 1 Criteria of properties for a good probiotic strain

This list should give a comprehensive guideline how to select single bacterial strains for probiotic use. However this list was updated and besides of human origin also dairy products and breast milk are used to isolate potential probiotic strains (Fontana et al.; 2013). To achieve a successful GRAS status it mustn't be pathogenic or toxic. To survive and grow in the in vivo conditions (human, animals), the probiotic strain must tolerate low pH and high concentration of conjugated and unconjugated bile acids. Of course the selected strain must be tolerated by the host even to people with reduced immune tolerance (Collins et al.; 1998). The strain should show good adhesive qualities for sufficient colonization in the human gut. In the final product the number of viable cells should be enough to confer the proposed health benefits. Further the selected strain should be compatible with the product matrix and desired characteristics during storage conditions should be maintained. (Fontana et al.; 2013)

1.4.1 *Lactobacillus johnsonii*

Lactobacillus and *Bifidobacterium* species are the most popular for the production of probiotic products due to their convincing beneficial effects on human health and their possession of GRAS status. However, this can't be generalized because each subspecies of one family shows different outcomes of immune stimulation and tolerance. (Collins et al.; 1998)

In general, probiotics should release a good amount of **anti-inflammatory** such as IL-10, IL-6 and TGF- β and low in **pro-inflammatory** such as IL-12, IL-23 and TNF- α cytokines. Moreover good adherence properties are necessary for colonization and hence achieve beneficial health outcome.

Marcinkiewicz et al. compared 3 different *Lactobacillus* strains (*L.reuteri*, *L.animalis/murinus* and *L.johnsonii*) in mice in terms of the production of cytokines. It was shown that *L.johnsonii* had higher IL-10 production than *L.reuteri* and lower than *L.animalis/murinus*. In IL-6 it was the other way round. However, in TNF- α *L.johnsonii* showed much lower production than *L.reuteri* (Marcinkiewicz et al.; 2007). A study with monoassociated *L.johnsonii* mice showed an increase in IgA+B-lymphocytes compared to germ-free mice 30 days after inoculation (Ibnou-Zekri et al.; 2003). However, they didn't give a comparison to conventionalized mice! Recently, Schiestl lab and others came to a similar result. A cancer sensitive mouse model (*Atm*^{-/-}) with defined intestinal microbiota composition was investigated to perform inflammation and genotoxicity testing. They came to the result that *L.johnsonii* treatment significantly reduced levels of the pro-inflammatory cytokines IL-1 β and IFN- γ , and elevated the levels of the anti-inflammatory cytokines TGF- β and IL-10.

Further, they have shown that *L. johnsonii* can **reduce systemic genotoxicity**. A significant reduction of natural killer cells and T-lymphocytes reflected by DNA damage in liver, spleen and blood was noticed compared to the control group with no *L. johnsonii* administered. (Yamamoto et al.; 2013)

Previous results on **adherence properties** showed that *L.johnsonii* strains (LA-1) have high adherence to intraepithelial cells (HT29 cells line), generally higher than that of the probiotic control strain *Lactobacillus rhamnosus* GG. (Vizoso Pinto et al.; 2007) To the same result came Zhang et al., the research group compared 6 selected *Lactobacillus* strains (*L.johnsonii* F0421, *L.acidophilus* IN3432 and IN3821, *L.paracasei* IN3623, *L.rhamnosus* IN4024 and 4025) on adherence properties. Between all 6 strains *L.johnsonii* showed the highest percentage of adhesion to HT-29 cells.(Zhang et al.; 2012)

In the same study they tested the same 6 *Lactobacillus* strains to their viability to gastric juice (pH 2 for 1h) and pancreatin solution (pH 8 for 4h incl. bile salt). *L. johnsonii* showed the highest survival rate to gastric juice treatment within all and is among the best in pancreatin solution intervention. (Zhang et al.; 2012)

Besides positive immunomodulatory effects and adherence properties, probiotics have another advantage as they **stimulate antimicrobial activities** against intestinal pathogens, directly. Many strains with high adhesion ability also showed high autoaggregation ability. *L. johnsonii* strains also coaggregated well with the intestinal pathogens *Listeria monocytogenes*, *Staphylococcus aureus*, *Escherichia coli*, and *Salmonella enterica serovar Typhimurium* (Vizoso Pinto et al.; 2007). Zhang et al. could find competitive adherence properties to entero-invasive *S. soneii* at HT-29 cells with a reduction of up to 48% depending on the various set-ups of the experiment (Zhang et al.; 2012). Furthermore, by using *L. johnsonii* as probiotic strain eradication of *Helicobacter pylori* infection was shown (Hsieh et al.; 2012).

1.5 Mouse models

To study the dynamic, ecologically diverse community of microbes that reside in the human gastro intestinal tract and to help us understand the biological complexities of the processes that govern host-microbiota symbiosis, various models were established. Those are *in-vitro* on e.g. epithelial cells like HT-29 or *in-vivo* mouse models like germ-free (GF), mono- or bi-associated, poly-associated or human flora- associated. Single strains are used to detect unique roles for their beneficial or detrimental effects in health and disease (Sekirov et al.; 2010). Within those groups various knock-out mice are available. Which animal model is used is dependent on the specific case or area of research and has to be discussed in detail. Each model of course has advantages and limitations and will be discussed shortly.

GF models provide an excellent base for research, to elucidate the mechanism behind inflammation response, signaling pathways and genotoxicity of every single strain in controlled environment and outstanding results have been

published. Together with genetically modified organism, GF studies have a high potential to provide new information in metabolic activity. However, gnotobiotics have some limitations. It is well known that gut microbiota is crucial for the proper development of the host especially the immune system. GF mice might not reflect what actually occurs in the natural composition as they were raised without microbiota. Hence it is difficult to transfer the results obtained in a germ free system to a conventional host. (Falk et al.; 1998)

Compared to gnotobiology (=colonization of GF animals with selected gut microbes), **mono- or bi- associated** allow investigations of host-microbe interactions in a simplified environment from the very beginning. Whereas a mono- colonized model can only demonstrate host-bacteria interaction, a bi-colonized model can also show microbe-microbe competition. For bi-associated mice predictions can be made about their ecological niches. However, the limitations are similar to the GF model. (Sekirov et al.; 2010)

Historically, the **poly-associated** model was developed by Russell W. Schaedler with eight defined bacterial strains to achieve a standardized gut microbiota for powerful research. A slightly revised model of this standardized poly-colonized model is used up to now. Because of different microbiota composition in conventional mice in various animal facilities even between cages, a standardized model makes it much easier to compare studies and housing confounders can be eliminated (Dewhirst et al.; 1999). To guarantee a high quality and accurate composition regularly tests (16S rRNA sequencing) have to be administered. This model represents dominant phyla like in a normal host, however the dynamics of a normal microbial- host and microbe-microbe cannot be demonstrated completely. (Sekirov et al.; 2010)

In **human flora- associated** (HFA) animals, ex-GF mice are inoculated with human fecal suspension. So far, it is not clear if HFA mice behave like conventional ones. This model is suitable to study dietary changes and therapeutic treatment such as probiotics and antibiotics and their impact to host ecology and metabolism. Further it makes an advantage to eliminate human differences in genetic and environmental factors by using a population of mice

with the same conditions in housing, diet and identical genetics. Besides, it can be used where the ethical commission would not allow treatments in humans like toxins, chemicals or carcinogens. (Hirayama et al.; 2005) Transferring human commensal microbiota to a mouse will not necessarily reflect a functionally identical equivalent of the original host intestinal environment. Furthermore there is no guarantee that the imported microbial mixture will be stable in HFA animals. (Sekirov et al.; 2010)

2 Methods and Material

2.1 Animals and husbandry conditions

For experiments in which I could participate at the Schiestl lab, different mice models were used. Wildtype mice (C57BL6/J) and *Atm*^{+/-} (heterozygote) both originally from Jackson Laboratory (Bar Harbor, ME) were housed and bred under standard conditions according to the Animal Research Committee at UCLA Department of Laboratory and Animal Medicine (DLAM). Mice were housed under 2 types of specific pathogen-free (SPF) conditions, where either sterile or non-sterile food, water and bedding were used (Yamamoto et al.; 2013). Additionally, these two colonies of mice harboring distinct microbial communities in their intestinal tract: conventional microbiota (CM) – refers to specific-pathogen-free - and restricted microbiota (RM) – refers to limited composition of intestinal microbiota. The model was created by Fujiwara et al. for colitis susceptibility. The exact composition can be seen elsewhere (Fujiwara et al.; 2008).

Mice either undergo radiation treatment with Si Ions (1,5Gy; for 5min 30sec; energy: 850MeV; 50 let) or high energy protons once. Also non- radiation treatment mice were used for some experiments.

2.2 Cultivation of *Lactobacillus johnsonii* and inoculation of mice

Lactobacillus johnsonii (LJ-RS-1) a mucosa-associated bacterium was found as one candidate by high-throughput 16S rRNA sequence analysis between two different mice colonies with different intestinal microbiota and various housing

conditions to have the ability to reduce systemic inflammation and genotoxicity. (Yamamoto et al.; 2013)

2.2.1 Growing *Lactobacillus johnsonii*

Lactobacillus johnsonii is inoculated from frozen glycerol stock onto Lactobacillus Selection Agar (LBS Agar) and incubated for two days at 37°C in an anaerobic chamber. Bacterial growth was collected with a sterile loop and washed with 1X phosphate buffered saline (PBS) solution for 3 times by centrifugation (3000 rpm for 10min) to wash out the nutrients from the agar and resuspended in 1X PBS. A final bacterial solution was adjusted to a density of 10⁹ CFU/50µl as determined by OD (0.8 for our settings) readings of serial dilutions at 600nm. The dilution scheme could be like the following (depending on the amount of bacteria collected from the Agar plates): 1:5, 1:5 + 100µl. The readymade suspension is kept at 4°C until inoculation on the next day.

2.2.2 Inoculating the mice

The mice are inoculated with 50µl of the prepared bacteria suspension. The gavage needle is kept upside down gliding with its own weight into the mouth. Due to the vulnerability of the mice the needle is not pressed into the mouth and not more than 50µl are inoculated at once not to drown the mouse. Fecal samples are taken before the first inoculation and after the treatment.

2.2.3 Reagents for bacterial growing and inoculation

Reagents	Company	Product no./ Code
Anaerobic Pack - Anaero	MGC Mitsubishi Gas Chemical Co, Inc.	2083LJ-3
BBL LBS Agar	Beckon, Dickinson and Company	211327
Dry Anaerobic Indicator strips	Beckon, Dickinson and Company	271051

Table 2 Reagents for bacterial growing and inoculation

2.2.4 Material for bacterial growing and inoculation

Material	Company	Product no./ Code
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Anaerobic Pack - Anaero	MGC Mitsubishi Gas Chemical Co, Inc.	2083LJ-3
BBL LBS Agar	Beckon, Dickinson and Company	211327
Dry Anaerobic Indicator strips	Beckon, Dickinson and Company	271051

Table 3 Material for bacterial growing and inoculation

2.3 Isolation of Intraepithelial Lymphocytes from mice

In order to see alterations among the local immune system it is necessary to harvest intestinal tissue and analyze lymphocytes from the gut epithelium. A slightly adapted protocol from Montufar-Solis et al. was used (Montufar-Solis et al.; 2006). The followed Percol separation was not established in the Schiestl lab so far. I started investigations for further applications.

2.3.1 Protocol

- 1) After collecting all necessary samples (blood, fecal) and data (body weight) euthanize the mouse with isoflurane.
- 2) Wait for approximately 3 – 5min and feel if the heartbeat stopped.
- 3) Fix the mouse on a Styrofoam plate, spray ethanol and open the stomach.



Figure 7 harvesting small intestine and colon for extracting intraepithelial lymphocytes

- 4) Extract large and small intestine (cecum goes with the large intestine) and place each in a separate labeled Petri dishes, each containing 10 ml

of ice-cold D10HS (Dulbecco's modified eagle medium –DMEM with 10% Horse Serum), and keep on ice until the next step. (Fig. 7)

- 5) Using a Petri dish on ice containing ice-cold D10HS, remove the Peyers Patches with forceps (they look like little white pimples on the intestines) as well as any attached vasculature.
- 6) Cut open the intestinal segments longitudinally and perform 3 serial washes with ice colds DMEM.
- 7) Cut tissue into ~1 cm segments and then place the segments into a 50 ml conical Falcon® tube containing 25 ml room temperature DMEM-DTT.
- 8) Lay conical tubes flat and incubate 20-30 minutes at 37°C in an incubator, shaking at 220 rpm.
- 9) Decant the supernatant into a new 50 ml tube containing a 70-µm filter (Fisher Brand Cell Strainers, 22-363-548).
- 10) Add 20 ml of room temperature DMEM to the tube with the tissue.
- 11) Vortex this tube for 30 seconds on high, and pour this supernatant through the same 70-µm filter into the same 50 ml conical tube.
- 12) Pellet supernatants at 1150 x g (Beckman J6M rotor JS-4.2) for 10 minutes at 4°C.
- 13) Resuspend cells in 5 ml of ice-cold DMEM and place on ice. These are the first-released IELs.
- 14) REUSE the first solution of DMEM-DTT to the tissue and repeat Steps 5-10.
- 15) Freeze IEL solution down to -80°C for further investigations or continue with Percoll separation.

Percoll Separation:

All solutions are prepared from a 90% isotonic Percoll solution (9 parts 100% Percoll and 1 part 10X PBS)

- 40% solution (ex. 4,4ml Percoll + 5,6ml DMEM 10%HS)
- 70% solution (ex. 7,7ml Percoll + 2,3ml DMEM 10%HS)

- 1) Pellet IELs (1500 rpm) for 10 minutes at 4 °C.
- 2) Resuspend the pellets in 3 ml 40% Percoll.
- 3) Place 4ml of 70% Percoll into the bottom of a 15 ml conical tube.
- 4) Gently overlay the 40% Percoll solution (containing the cells) onto the 70% Percoll.
- 5) Centrifuge at 1500 rpm for 30 minutes at RT.
- 6) Collect cells (400µl) at the interface and add this solution and wash by centrifugation (1500 rpm) in 5ml RPMI-1640.
- 7) Repeat step 3.-6.

2.3.2 Material for Isolation of Intraepithelial Lymphocytes from mice

Material	Company	Product no./Code
Centrifuge	Backman Coulter	Microfuge 18
DB Falcon® tube (50ml/15ml)	DB Bioscience	352070
Incubator Shaker	New Brunswick Scientific Co. Inc.	Series 25
Petri Dish Fisherbrand	Fisher Scientific	0875712
Strainer DB Falcon	DB Bioscience	352350
Vortex	Fisher Scientific	12-812

Table 4 Material for Isolation of Intraepithelial Lymphocytes from mice

2.3.3 Reagents for Isolation of Intraepithelial Lymphocytes from mice

Reagent	Company	Product no./Code
Dulbecos Modified Eagle Medium (DMEM) 1X	gibeco by life technologies	11960-044
Equine Serum	HyClone	AJG10637
Percoll ®	Sigma-Aldrich	P-1644
Pierce® - DTT (dithiothreitol)	Thermo Scientific	20291
RPMI-1640 1X	gibeco by life technologies	11875-093

Table 5 Reagents for Isolation of Intraepithelial Lymphocytes from mice

2.4 γ H2AX foci determination on Blood Lymphocytes

2.4.1 Background and principles of γ H2AX assay

DNA double strand breaks (DSBs) are serious lesions that can initiate genomic instability, ultimately leading to cancer (McKinnon et al.; 2007). Due to various endogenous and exogenous factors DNA damage can occur. These can be classified according to the underlying cause as followed: (a) exogenous: direct interaction with a damaging agent such as toxic chemicals or pharmaceuticals, radiation/UV, physical activity, tobacco smoke and nutrition; (b) endogenous: reactive oxygen species (ROS), metabolic processes, deficient repair, telomere erosion, inflammation, cellular respiration and programmed biological processes. (Bonner et al.; 2008; Bensimon et al.; 2011)

Human and animal cells have to deal with thousands of DNA lesions per day. Therefore a good working repair mechanism is from enormous importance to keep body functions ongoing. Of all DNA strand breaks, not all of them are accidentally. Endonucleases play an important role in crossing-over during mitosis and meiosis as well as programmed cell death leads to DNA lesions. (Rogakou et al.; 2000; Fillingham et al.; 2006)

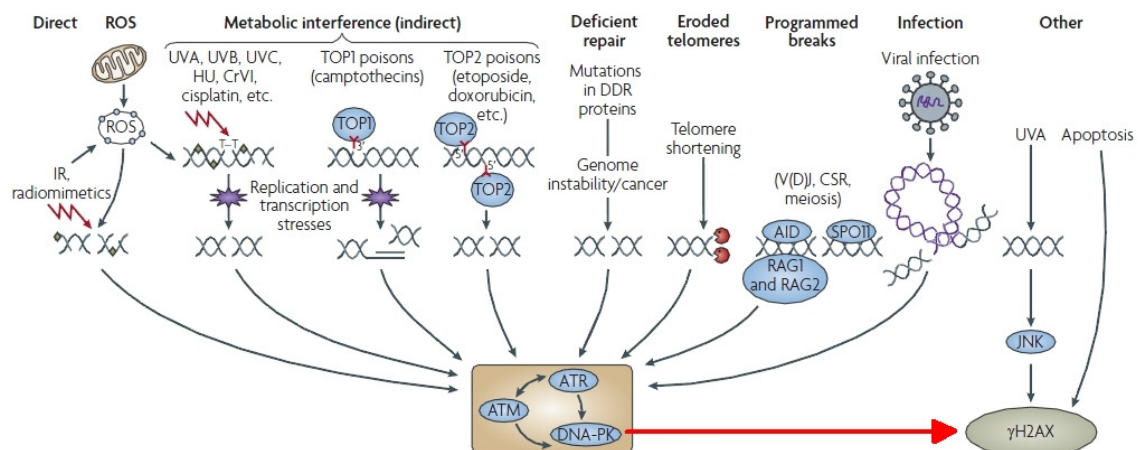


Figure 8 This illustration shows various categories of origins of DSBs and how they lead to γ H2AX phosphorylation by three kinases ATM, ATR and DNA-PK. Adapted from (Bonner et al.; 2008)

In eucariotic cells, DNA is packed into nucleosomes, which consists of winded DNA around proteins and are arranged in higher structures to form chromatin.

To have a closer look at one nucleosome it consists of 140 to 145 base pairs of DNA and eight histone proteins, two from each of the H2A, H2B, H3 and H4. To stabilize the structure a fifth histone protein H1 is responsible and is acting as a bridge between nucleosomes. The H2A family consists of 3 subfamilies H2A1- H2A2, the H2AZ, and the H2AX; in mammals the H2AZ represents about 10% of the H2A complement, the H2AX represents 2–20%, and the H2A1- H2A2 represents the balance. Each nucleosome contains two molecules of the H2A family. (Rogakou et al.; 1998)

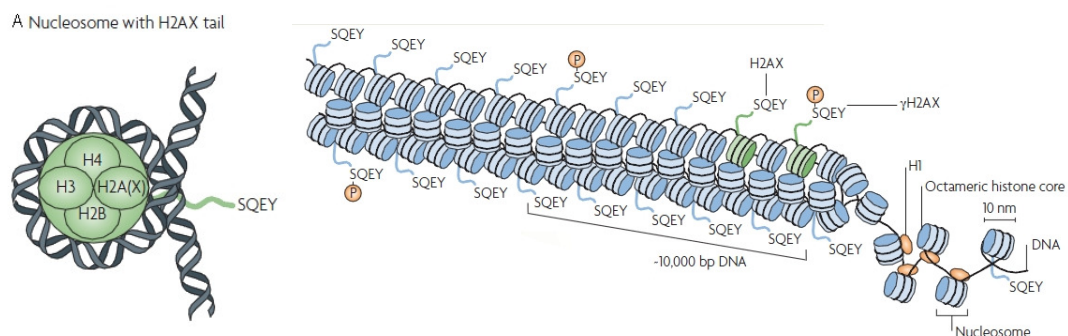


Figure 9 Left: H2AX is a component of the octamer (4x2) of histones packaging DNA into a nucleosome, while many nucleosomes form the chromatin. Right: The nucleosomes form a fibre containing H2AX molecules in every fifth nucleosome on average in mammals. Approximately 10% of the H2AX molecules are phosphorylated at any one time in a focus. (Bonner et al.; 2008)

H2AX omega 4-serine 139 becomes rapidly phosphorylated after DNA double strand breakage. Just one DNA double strand break leads to hundreds of copies of phosphorylated H2AX (γ H2AX), which form foci covering many megabases of chromatin and encompassing the DSB location. Immediately, DNA damage response proteins are recruited which in turn activate DNA repair processes. There are 3 main PI3K-like kinases involved ataxia telangiectasia (ATM), ataxia telangiectasia Rad3 related (ATR) and DNA- dependant protein kinase (DNA-PK) as well as lots of other checkpoint and DNA repair proteins (Fig. 8). Thus, H2AX represents a key factor in the repair process of damaged DNA. (Dickey et al.; 2009) Immediately after DSBs happen γ H2AX formation begins. Approximately between 9 and 30 minutes after DSBs occur, large numbers of γ H2AX molecules accumulate to form foci (Rogakou et al.; 1998).

Measurement of γ H2AX foci by fluorescent staining has become a popular method for detecting DSBs as the foci are easy to identify with antibodies and H2AX phosphorylation is a highly sensitive biological marker of DSB formation. Latest makes it a good marker for improvement of therapeutic and pharmacological interventions (Barber et al.; 2007). The detection and quantification involves a two step detection, first a γ H2AX primary antibody and a fluorescent antibody. The detection requires a immunofluorescent microscope (Kuo et al.; 2008).

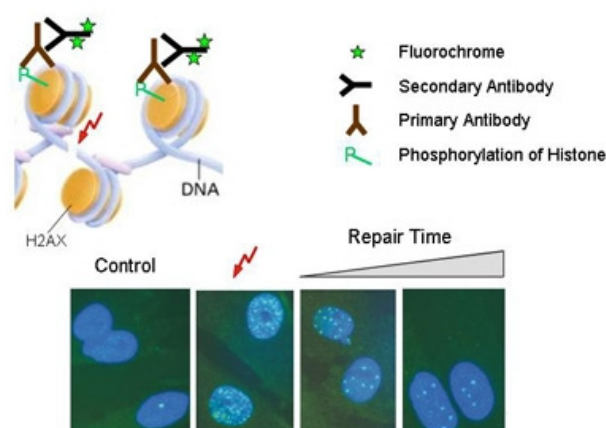


Figure 10 Principle of γ H2AX assay and foci demonstrating fluorescent pictures (<http://www.auntminnie.com/index.aspx?sec=ser&sub=def&pag=dis&ItemID=85566>)

Recently, it has been shown that the gammaH2AX assay is a hundred times more sensitive than the similar and widely used comet assay (Ismail et al.; 2007). Thus, it makes the γ H2AX assay one of the most used assays for DNA double strand breaks also because the preparation is simple and easy. The major field of usage is to detect DSBs after irradiation, however the work performed in Schiestl lab was different. The goal was to see long term effects after irradiation because of different gut microbiota and husbandry conditions. The hypothesis is to see a reduction of DNA damage of single bacterial strains reflecting a decrease in inflammation.

Limitations of the γ H2AX assay (Löbrich et al.; 2010):

- Senescent cells show foci at eroded telomeres without DNA damage

- Mitotic chromosomal breakage are visualized, approximately every 1 out of 10 foci DSBs is because of mitotic cell activity
- Cell cycle activity in S-Phase can lead to increased foci formation
- A decreased sensitivity can be estimated in cell lines with a lack in ATM or ATM-dependant signaling proteins
- It is commonly recommended to score DNA DSB 30min after irradiation exposure, however rapid repair mechanism could underestimate the amount of foci
- A high induction of single strand breaks does not necessarily mean high DSBs (e.g. 0.1mM H₂O₂ cause high SSB detected by comet assay however only a few γ H2AX foci).

2.4.2 Protocol for γ H2AX assay

Preparing coverslips to drop cells

- Make sure coverslips have been cleaned: 2 hours in concentrated HCl, washed in water overnight. Store cleaned coverslips in 100% Ethanol.
- Flame the coverslips before use to remove Ethanol.
- Add 150 μ l of 100 μ g/ml poly-D-lysine (in H₂O) to each coverslip. Swish the coverslip so that the poly-D-lysine covers it all. Incubate for 5 min at room temperature (RT), aspirate the solution.

Dropping the cells

- Incubate 50 μ l blood in 250 μ l Erythrocyte lysis buffer (Qiagen). Leave on ice for 15-30 min, vortex a few times. Centrifuge for 7 min at 2500 RPM.
- Remove supernatant, resuspend cell pellet in 150 μ l Erythrocyte lysis buffer, incubate for 1 min, centrifuge for 7 min at 2500 RPM, resuspend pellet in 50 μ l PBS.
- Drop 50 μ l cell suspension onto poly-D-lysine coated coverslips. Pipette gently cells onto coverslip so that they are covering all the coverslip.

- Add 1 ml of PBS to prevent cells from drying out. Put coverslips in a 8-well Dish Nuclon Delta Treated (NUNC #167064, www.nuncbrand.com). Put one coverslip per well.

Fixing and permeabilisation

- Dump out PBS and fix cells by adding 2 ml of 4% paraformaldehyde in PBS. Incubate for 10 min at RT.
- Aspirate paraformaldehyde
- Wash once with PBS by adding 2 ml of PBS and incubating for 3 min at RT.
- Remove PBS and add 2 ml 0.5% TritonX-100 in PBS for 10 min (time is crucial).
- Wash with 2 ml of PBS 5 times (add 2 ml and dump out, in total 5 times).

Blocking

- Remove PBS and add 2 ml of 10% Equine Serum (ES) in PBS.
- Incubate overnight at +4 °C.

Primary Antibody

- Prepare the primary antibody in 10% ES in PBS (1:400)
- Add 50 µl of primary antibody solution on each cover slip in humidity chamber, put squire parafilm onto coverslips to avoid drying, cover the chamber to prevent from drying out.
- Incubate for 1.5 h at RT.

Second Blocking

- After primary antibody incubation is complete, wash the coverslips 3 times with 2 ml 0.1% TritonX-100 in PBS for 5 min each time.
- Add 2 ml of 10% ES in PBS.
- Incubate for 1 h at RT.

Secondary Antibody

- Prepare the secondary antibody in 10% ES in PBS (1:150)
- Add 50 µl of secondary antibody solution on each cover slip in humidity chamber; put square parafilm onto coverlips to avoid drying.
- Incubate for 45 min at 37°C. Cover the plate with aluminum foil to avoid exposure to light.
- Wash the coverslips 3 times with 2 ml 0.1% TritonX-100 in PBS for 5 min each time.

Preparing Slides

- Put 5 µl of DAPI solution (1 volume of DAPI in Vectashield : 5 volumes of Vectashield) onto a pre-labeled slide.
- Pick up coverslip (remember what side the cells are on), remove excess liquid using edges of a paper towel. Place the coverslip cells facing down onto DAPI mounting media.
- Analyze γ-H2AX foci under 100x or store slides at -20°C to avoid exposure to light

2.4.3 Material for γH2AX assay

Material	Company	Product no./ Code
8-well Dish Nuclon Delta Treated	Thermo Scientific	nunc#167064
Centrifuge	Backman Coulter	Microfuge 18
Coverslips		
Microscope	Olympus	OlympusBX51
Microscope slides	Fisher Scientific	12-550-15
Parafilm	Pechiney-Plastic Packaging	
Pipettes	Pipetman P1000/P200	
	Rainin SL 20	
Shaker	VariMix	
Vortex	Fisher Scientific	12-812

Table 6 Material for γH2AX assay

2.4.4 Reagents for γ H2AX assay

Reagent	Company	Product no./Code
<u>Antibody (first):</u> Anti-phospho-H2A.X (Ser139) Polyclonal Antibody IgG	Millipore	Cat.# 07-164
<u>Antibody (second):</u> Flourescein (FITC)-conjugated AffiniPure F(ab') ₂ Fragment Goat Anti-Rabbit IgG, F(ab') ₂ Fragment Specific (minimal cross-reaction to Human Serum Proteins)	Jackson ImmunoResearch Laboratories, Inc.	111-096-047
Equine Serum	HyClone	AJG10637
Erythrocyte Lysis Buffer	Qiagen Science	1014617
Ethanol	Fisher Scientific	128173
HCl	Fisher Scientific	CAS 7647-01-0
Paraformaldehyde 4% in PBS	Affymetrix	19943
PBS – Buffer	Apex	12135JK
Poly-D-lysine	Sigma-Aldrich	1001297502
Triton X-100	Promega	0000017643
Vectashield	Vector Laboratories	H-1400
Vectashield with DAPI	Vector Laboratories	H-1200

Table 7 Reagents for γ H2AX assay

2.5 Micronucleus determination in Blood Erythrocytes

2.5.1 Background and principles of micronucleus formation

The micronucleus (MN) assay is a recently used and well established method to measure genotoxicity in form of chromosomal damage and is widely used, both in vivo and in vitro. It is commonly used to assess chemical substances on genetic mechanisms, pharmacokinetics, and DNA-repair process for acute and chronic effects. Of the same importance are studies that show adverse effects especially in epidemiological studies. (Krishna et al.; 2000)

Historically, the micronucleus assay was performed on bone marrow because erythropoiesis with proliferation and maturation stages takes place in bone marrow and spleen. Later it was established also for peripheral blood erythrocytes after they found histological liver samples from mice which show micronucleus in blood vessels. Nowadays it is mainly used for mammalian erythrocytes and human lymphocytes. (Heddle et al.; 2011)

Erythropoiesis and formation of micronuclei:

It is now well-known that MN mainly originate during anaphase from lagging acentric chromosome or chromatid fragments which are caused by misrepair of DNA breaks or unrepaired DNA breaks, but this is only likely if the damage load exceeds the repair capacity. Further they can occur during anaphase due to malsegregation of whole chromosomes which cannot properly attach to the spindle, defective checkpoint genes and defects of in kinetochore proteins. All kinds of damage are associated with the development and progression of tumors. (Fenech et al.; 2011)

The process of erythropoiesis is shown in the following figure.

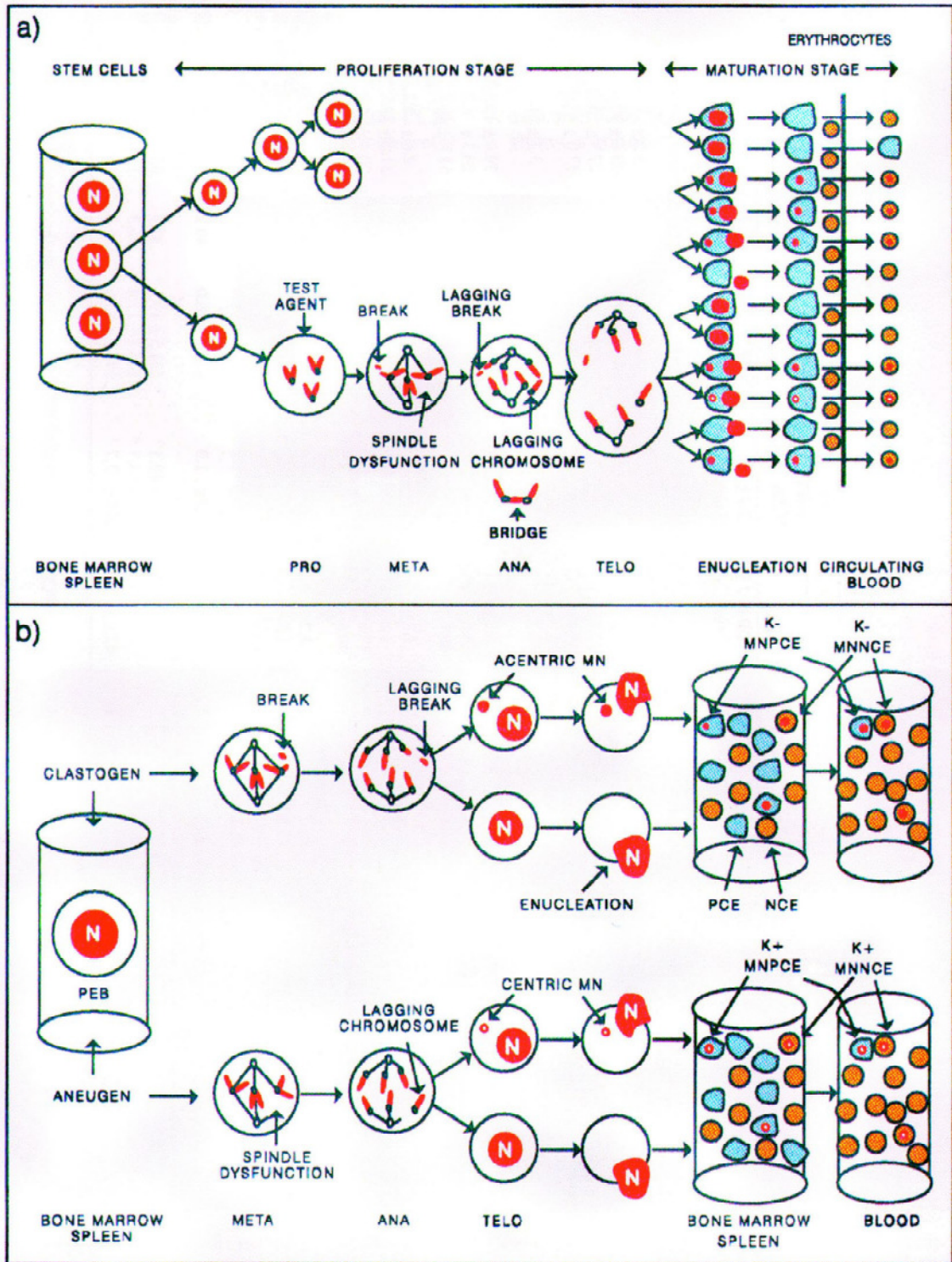


Figure 11 Erythropoiesis (a) The process of erythropoiesis in vivo; (b) the mechanism of micronucleus formation in the polychromatic erythrocytes (PCEs) and normochromic erythrocytes (NCEs). (Krishna et al.; 2000)

During erythropoiesis stem cells from hemopoietic organs develop erythrocytes through a proliferation and maturation stage. In the proliferation stage cells

continue to divide and in this sensitive phase administered testing chemicals may cause MN due to the reasons mentioned above. These anomalies like chromosome fragments or whole chromosomes may form micronuclei in cytoplasm. A regular cell division or a proper break down of the nuclei to daughter cells is defective. In the next stage, during maturation, when an erythroblast develops into a polychromatic erythrocytes (PCE), the main nucleus is unfolded and any micronucleus that may has been formed may remain behind in the otherwise enucleated cytoplasm. This is the main reason why a visualization of micronucleus can be performed, because they lack a main nucleus. With time, the polychromatic erythrocytes lose RNA and become normochromatic erythrocytes (NCE) containing primarily hemoglobin. Later in the maturation stage, mostly NCE move into the peripheral blood compartment. (Krishna et al.; 2000)

In this work cells were stained with Wright's Giemsa (protocol see below). This method is very easy to perform, whereas it takes a long time to count cells under the microscope. For each sample two microscope slides were prepared and 4000 cells were counted on each slide. It is suggested that at least a 1000 samples should be counted (Krishna et al.; 2000).

See pictures below for examples considering exclusion criteria as followed:

- The micronucleus is in focus when the cell is in focus to avoid mistakes by blue colored crystals of the staining solution
- Appears black under the microscope itself and dark or intensive purple on the screen using the camera
- I always started with a micronuclei on screen to have a reference (color)

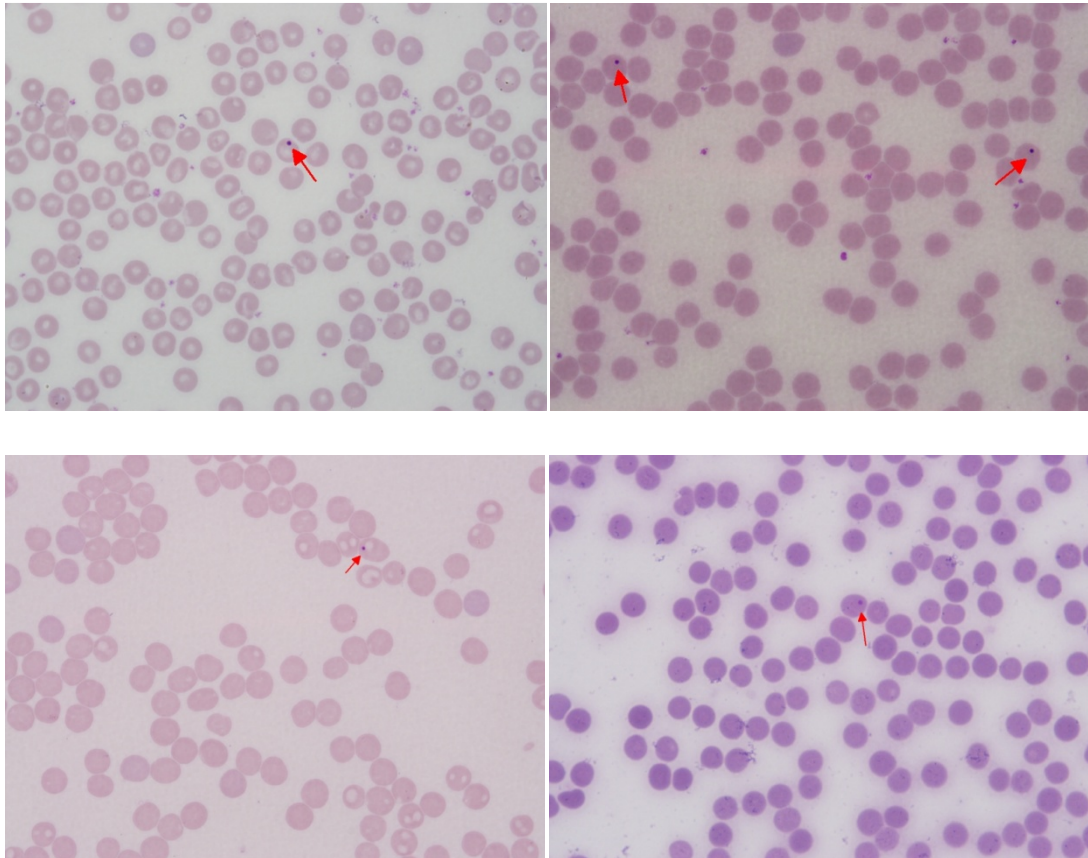


Figure 12 Those four pictures show micronuclei under the microscope (100x) with Wright's Giemsa staining

2.5.2 Protocol for micronucleus determination

- Pipette 3µl of fresh blood onto a microscope slide and smear
- Put the slides in methanol for two minutes and let them dry overnight
- Put slides in Accustain® Wright's Giemsa for five minutes
- Wash slides two times in water
- Analyze Erythrocytes under 100x and store at RT

2.5.3 Material for micronucleus determination

Material	Company	Product no./ Code
Microscope	Olympus	Olympus BX51
Microscope slides	Fisher Scientific	12-550-15
Pipettes	Rainin SL 20	

Table 8 Material for micronucleus determination

2.5.4 Reagents for micronucleus determination

Reagent	Company	Product no./ Code
Accustain® Wright-Giemsa stain	Sigma-Aldrich	5K200R4

Table 9 Reagents for micronucleus determination

2.6 Single cell gel electrophoresis (comet assay) for lymphocytes from whole blood

2.6.1 Background and general principles of the alkaline comet assay

The comet assay is together with the micronucleus assay and γ H2AX one of the most widely used assay to detect DNA damage. Historically, start up procedures to measure DNA strand breaks were mentioned already in the 1970th. In 1984 Ostling and Johanson first described a method under neutral conditions to measure only DNA double strand breaks due to relaxation of DNA supercoils. Later in 1984 it was first developed under alkaline conditions and it was possible to assess both, single and double strand breaks (Fairbairn et al.; 1995). From this time on a yearly increasing number of papers based on this method dealing with DNA damage and repair were published. Nowadays, the comet assay is well established and an OECD approved method for genetic toxicity testing e.g. pharmaceutical, radiation experiment, environmental hazards etc. The method is easy to perform and economic, however there are some limitations and variations within the protocol which makes it difficult to compare between similar papers (Tice et al.; 2000).

This assay mainly neutral and alkali and its modified versions (application of enzymes, electrophoresis, time, voltage,...) can be used with nearly all different kinds of eukaryotic cells and origin from bacteria, plants, algae, animals to humans. This is described in more details elsewhere. (Dhawan et al.; 2009)

The general principle (Fig.13) of the comet assay is the migration in the electric field of negative charged DNA to the anode. Therefore single cells – in this case mouse lymphocytes from whole blood – are embedded on agarose gels and lysed with detergent and high salt to form a nucleoid of supercoiled DNA loops.

DNA breaks relax those supercoils and form a so-called “halo”. The more strand breaks the slower DNA can move to the positive charged anode and the amount of DNA in the tail represents proportionally the amount of strand breaks, respectively. (Collins; 2004)

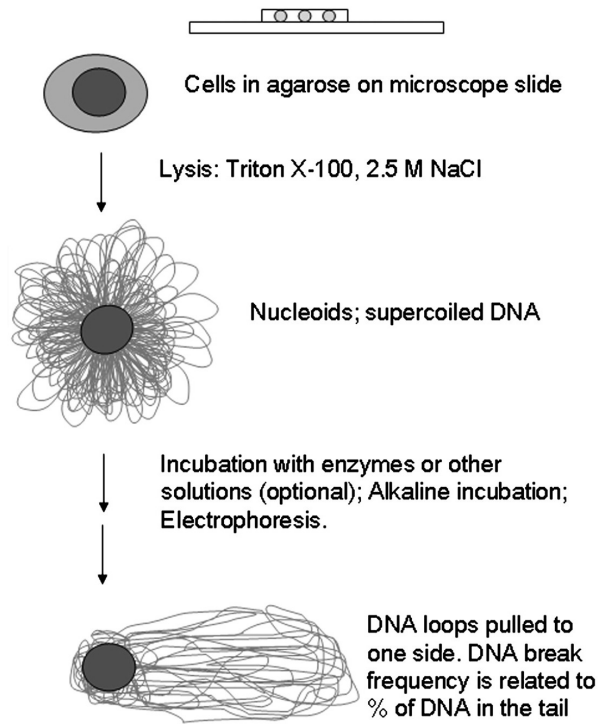


Figure 13 General principle of comet assay
<http://mutage.oxfordjournals.org/content/24/5/383/F1.expansion.html>

The comet assay is named after the appearance receiving from the microscopic detection after fluorescent staining. Whereas the head contains the intact DNA and the tail consists of DNA fragments. (Olive et al.; 2006)

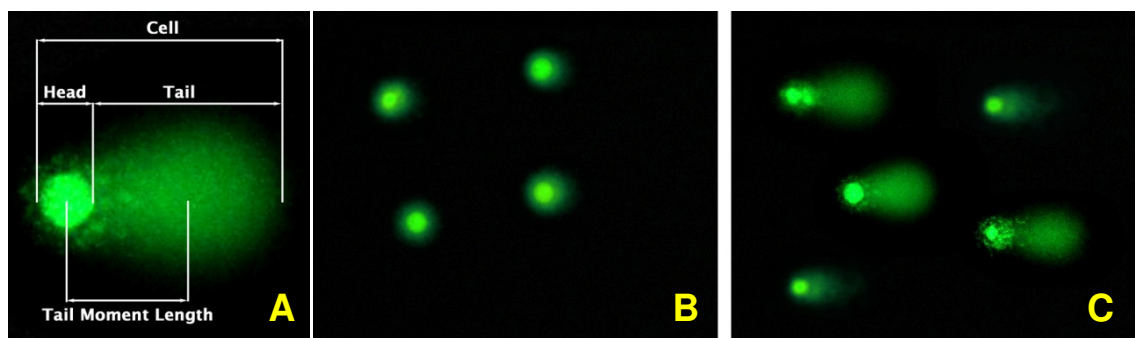


Figure 14 Fluorescent pictures of comets under the microscope. A) Description of comet components. B) untreated sample non-/low- DNA damage C) high DNA damage rate. (<http://www.cellbiolabs.com/comet-assay-kits-96-well>)

The pictures above show visualized fluorescence pictures from single cells. Each green dot represents one cell. In B very low damage is visible whereas C shows medium to high DNA damage.

Advantages of the comet assay are the following (Olive et al.; 2006; Azqueta et al.; 2013):

- Low costs,
- Fast and easy procedure,
- Results can be obtained on the same day,
- High sensitivity for measuring low levels of DNA damage, both single and double strand breaks in alkaline comet assay (which was used)
- Small sample size from 10.000 to 50.000 single of various cell types and origin
- Fresh and frozen (rapidly to minus 80 °C) samples can be used
- Flexibility to use proliferating as well as non-proliferating cells
- Generated data allow robust types of statistical analysis
- The ability to analyze single cells that might respond different to other identifying subpopulations

However there are some limitations which should be mentioned (Fairbairn et al.; 1995; Olive et al.; 2006; Azqueta et al.; 2013):

- Slight differences in protocols and technical variability (electrophoresis, time, agarose concentration, buffer...)
- Requirement for a viable single-cell suspension (intact cells, if samples contain too many necrotic and apoptotic cells, accurate results cannot be obtained)
- No information on fragment size
- Aneugenic effects and epigenetic mechanism are not detected
- Single cell data might be rate limited
- Small cell sample leading to sample bias
- Various background level (age, gender, stress, inflammation...)
- Scoring comets is tedious (eye visualization, program)

- Cross-links (from e.g. chemicals) will block detection

2.6.2 Methodology – step by step

As written above many different versions are found in literature. A complete standardization of parameters is not achievable, however to compare results between different laboratories, it is necessary to have a closer look to the factors that affect the performance of this assay.

1) Slide preparation

For preparing the agarose gel different version were found in literature (Hartmann et al.; 2003). In this work a 2 layer version was used. First a 1% standard agarose gel on a GelBond film was prepared. The diluted samples were mixed into a 1% low melting point agarose (LMP) which, after drying the standard agarose, were put on imprinted circles.

The density of the agarose can affect the extent of DNA migration in the electric field. A concentration of 0.5 to 1% LMP agarose gel is commonly reported. An equally important impact of the quality of the comet assay results from the amount of cells. High cell densities can lead to overlapping of comets. (Azqueta et al.; 2013)

2) Lysis

The reported lysis times vary from a minimum of 30 minutes to overnight. Citation missing! Azqueta et al published an optimum of 40 minutes (Azqueta et al.; 2013), whereas Hartmann et al talk about a minimum of 1h (Hartmann et al.; 2003) both in alkaline conditions (>13 pH). The lysis solution consists of detergents (Triton X-100) and high salts (2.5M NaCl) concentration. DMSO acts as radical scavenger if whole blood is used. During lysis histones are solubilized by high salt, nucleosomes are disrupted; cytoplasm, membranes and nucleoplasm are removed. (Collins; 2004) However the negative supercoiling of the DNA survives as long as the DNA is intact.

3) Alkaline treatment

After 1h in the lysis solution an alkaline treatment (pH 13) was followed. Prior to the electrophoresis an unwinding time of 20min (depending on the cell type) is considered as being enough to relax the supercoiling structure. The more DNA strand breaks occur the more DNA loops will be relaxed and the more DNA is detectable in the comet tail. (Azqueta et al.; 2013) The agarose gel is already placed in the electrophoresis chamber and kept in the fridge at 4°C.

4) Electrophoresis

During electrophoresis DNA is migrating through an electric field towards the anode. As mentioned above the more DNA damage is present the bigger the comet appears. The electrophoresis is the part with the most possibilities for variations in time, voltage and currents. Experiments show a constant increase in DNA migration the higher the voltage (up to 1.48 V/cm) and as well a constant increase in % DNA in the tail by increasing the time (up to 40min). All variations were tested on lymphocytes. Azqueta et al. came to the following conclusion that 20min at 1.15V/cm and 30min at 0.83V/cm are considered the most reliable. An increase in current due to changing the volume of the electrophoresis solution implies a decrease in voltage and is also seen in a decrease of % DNA in the tail. (Azqueta et al.; 2013)

5) Neutralization

After electrophoresis, the alkalized agarose gel is neutralized by washing with neutralization buffer for 3 times each 5 minutes.

6) Staining and scoring

There are several possibilities to stain the comets such as ethidium bromide, 4,6-diamidino-2-phenylindole (DAPI), SYBR Green and SYBR Gold. Which magnification is most appropriate depends on the cell types. For measuring comets various methods are available from visual scoring to computer based automatic scoring. Both mentioned have advantages and disadvantages.

2.6.3 Solution preparation for comet assay

Lysis solution

- 2.5M NaCl
- 0.1M EDTA pH 10
- 10mM Tris
- 1% Triton X-100 (add immediately before use)
- 10% DMSO (add before use,)

Electrophoresis solution (pH 13)

- 0.3M NaOH
- 1mM EDTA (from stock solution: 200mM EDTA pH 10)

Neutralizing buffer

- 0.4M Tris
- Conc. HCl to pH 7.5

H₂O₂ stock solution

- Hydrogen peroxide solution (30%)
- deionized H₂O

11.5 µl of the hydrogen peroxide solution was mixed with 1 ml deionized H₂O to receive a 0.1M stock. Before usage dilute 15µl of the 0.1M stock solution in 30ml PBS (50µM).

Normal melting agarose

- 1% standard agarose
- 1X PBS

40ml are enough to cover the GelBond film.

Low melting point agarose (LMP)

- 1% LMP

- 1X PBS

Prepare 1ml aliquots for storage (-20°C).

2.6.4 Protocol for comet assay

The comet assay was done as described previously with some small adoptions (Singh et al.; 1988):

Slide preparation

- Coat GelBond film with 1% standard agarose in PBS and pour hot agarose on the film and make sure it is completely covered. Let it solidify at room temperature for 5-10 min.
- Imprint rings (~1 cm diameter) on the surface of the gel using a clean glass tube.

Blood collection and preparation

- Collect blood in a tube containing Na₂EDTA (1 vol. Na₂EDTA + 9 vol. whole blood) and mix it up by inverting tube up and down.
- Add 10µl blood into 110µl PBS 1X
- Keep samples on ice

Embedding cells in agarose

- Prepare 1% low melting point (LMP) agarose in PBS
- For each sample prepare an EP tube containing 200 µl of 1% LMP agarose, incubate tubes in 37°C water bath
- Add 60µl of cell suspension (out of the 120µl) to 200 µl LMP agarose, mix well by pipetting up and down several times, put in 37°C in water bath for 1 min.
- Take 25 µl of cell suspension in LPM agarose and pipet into an imprinted circle on a precoated GelBond film (do in triplicate). Let it solidify for 5 min.

- To induce DNA damage as a control sample prepare a 0,01M H₂O₂ solution and add 10µl to the blood solution (10µl whole blood + 110µl PBS)

Lysis (1 h)

- Immerse Gelbond slides in chilled lysis solution (2.5 M NaCl, 10 mM Tris, 100 mM Na₂EDTA, pH 10.0, with 1% Triton X-100 and 10% DMSO added fresh) and incubate for 1 h at 4°C to remove cellular proteins and liberate DNA (cover with plastic foil).

Alkaline treatment (20 min)

- Transfer the slide to a horizontal electrophoresis chamber (BioRad, Hercules, CA) filled with fresh, chilled electrophoresis buffer (300 mM NaOH and 1 mM EDTA, pH 13) and leave for 20 min at 4°C to allow DNA unwinding.

Electrophoresis

- Perform electrophoresis in the same buffer at 300 mA (~20V depending on tank dimensions) for 40 min at 4°C. Adjust 300 mA by lowering or increasing the volume of the buffer (approx. volume 500 ml).

Neutralisation

- After electrophoresis wash the slide with neutralizing buffer (400 mM Tris-HCl, pH 7.5) by three times 5 min incubations at 4°C.

Staining

- Dispense 25 µl of SYBR Gold (1/10,000 dilution of stock solution from Molecular Probes, 495 nm excitation, 537 nm emission) on each circle of the slide.
- Incubate for 5 min at room temperature, rinse with distilled water to remove excess dye. Visualize comets under fluorescent microscope (FITC filter) under 10x or higher magnification.

Analysis

- For each sample three equal imprints were made and of each imprint ~ 10 pictures were taken (between 25 and 30 per sample)
- Pictures were uploaded to the casp.exe program (Comet Assay Software Project, <http://casp.sourceforge.net/>) and olive tail moments were measured (in total between 100 and 200 comets)

2.6.5 Material for comet assay

Material	Company	Product no./ Code
GelBond Film	Lonza	53748
Electrophoresis Chamber	Owl Scientific Inc.	Model #D-3
Microscope	Olympus	OlympusBX51
Power Supply	Thermo Electron Corporation	EC4000P

Table 10 Material for comet assay

2.6.6 Reagents for comet assay

Reagent	Company	Product no./Code
Agarose NuSieve 3:1	BioProducts	50090
Dimethylsulfoxide	Fisher Scientific	116070
HCl	Fisher Scientific	121507
H ₂ O ₂	Kroger	L0012462FA
Low Melting Point Agarose	invitrogen life technologies	15517-022
NaCl		
Na ₂ EDTA	Fisher Scientific	075032
NaOH	Fisher Scientific	107702
PBS – Buffer	Apex	12135JK
CYBR Gold	Molecular Probes	
Tris Base	Fisher Scientific	107702
Triton X-100	Promega	0000017643

Table 11 Reagents for comet assay

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