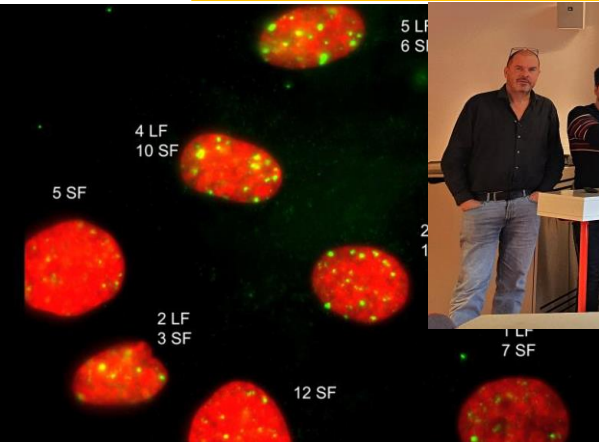
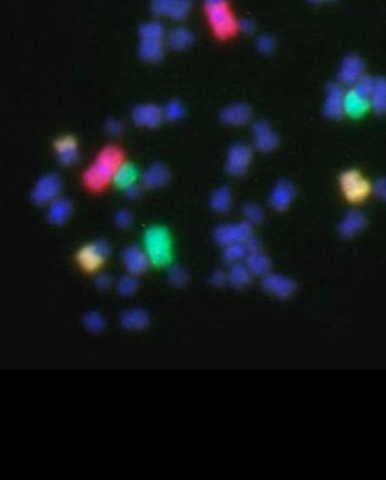
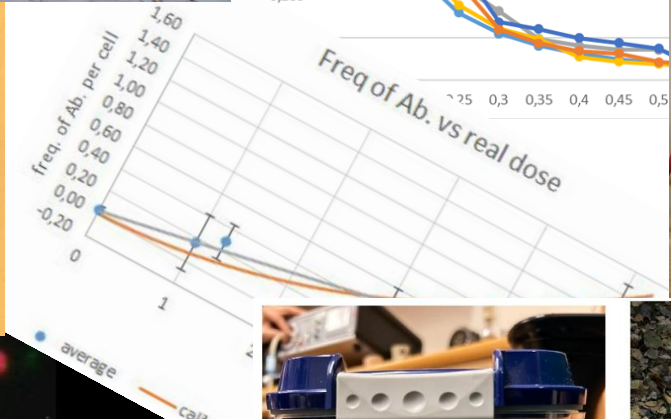
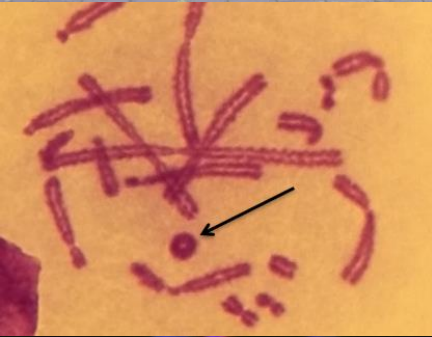
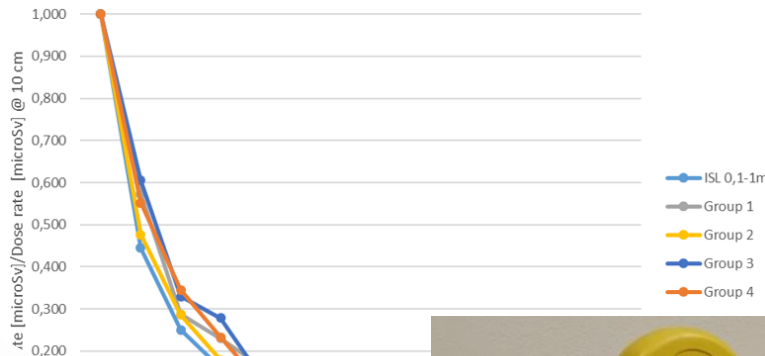


A RadoNorm short course entitled: CELET: Cellular and genotoxic effects of high and low LET ionising radiation – introduction to radiation biology

Stockholm University, Sweden
11.11.2024 – 22.11.2024



Runaway Train Law vs. Inverse Square law (Distance 0,1-1m)
Background subtracted



A RadoNorm short course entitled:

CELET: Cellular and genotoxic effects of high and low LET ionising radiation – introduction to radiation biology

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Version 1 – 2023 12 21

Aim of the course

The aim of the course is to acquaint students with techniques of studying genotoxic effects of ionising radiation which are of relevance for RadoNorm and the broad field of radiation research. The target group are students and young researchers with various backgrounds who want to get a basic introduction to biological effects of radiation. The course will contain both lectures and practical laboratory work. The lectures will focus on various aspects of biological effects of low and high LET ionising radiation as well as on techniques to detect them using cytogenetics and immunogenetics.

The practical part will focus on teaching laboratory techniques used to study genotoxic radiation effects on cells: 1) harvesting cells and analysing chromosomal aberrations and micronuclei by the Giemsa method; 2) in situ hybridization with whole chromosome probes and analysis of translocations; 3) gamma H2AX focus assay; 4) basics of high and low LET radiation detection and dosimetry.

The course will last 2 weeks and will be nested in a 10 week “introduction to radiobiology and cellular toxicology” course held at the Stockholm University (SU). The regular SU course is held for a maximum of 16 students. CELET will harbour a maximum of 12 students, bringing the total number of participants to 28.

Information for applicants: The course is open to any postgraduate student or researcher working in an EU academic institution. There is no course fee and every student will receive a financial support equivalent to 50 Euro per day to cover the costs of lodging. The University has no lodging possibilities so lodging will be offered in a youth hostel. Alternatively, the applicant can find her/his own lodging but the sum of financial support will remain 50 Euro per night. No other financial support will be provided.

People wishing to apply should submit by mail the following documents to Andrzej Wojcik at andrzej.wojcik@su.se:

1. A letter of application
2. A CV with a description of the scientific career
3. A supporting letter from the supervisor/head of laboratory

The **deadline for applications is September 30th 2024**. Information confirming the acceptance will be sent by 1st October 2024. A diploma, equivalent to 3 ECTS points, will be issued to each participant after the course.

Course description

To facilitate work in the lab the students will be divided into 4 groups. Each group will carry out one experiment with different endpoints. At the end of the course students will present and discuss their results.

The course lectures will be held in the morning hours during week one. Practical work will be carried out in the afternoon hours of week 1 and during the whole days of week 2. The practical work will be divided into a “lab-teaching” part and a “results-analysis” part. Each group will learn techniques 1-4 (see above). Hence, each student will spend 4 time-blocks in the lab, where she/he will carry out the steps associated with a technique. The rest of the practical time will be devoted to learning how to analyse cells on microscopic slides/images. The rationale for dividing the students into small groups is that it will allow them to really perform the work and not only watch a demonstration. Each group will be supervised by an experienced employee of the SU.

Each group will also learn how to irradiate cells using the exposure facilities at SU: low-dose rate ¹³⁷Cs exposure facility, high dose-rate ¹³⁷Cs exposure facility, X-ray facility and ²⁴¹Am alpha-exposure facility. Although students will learn how to generate microscopic slides/images that can be used for analysing the results, the scoring part of the course will be carried out using slides/images prepared beforehand by the SU employees. This strategy will guarantee high quality slides/images for scoring. At the end of the course the achieved results will be collated, statistically analysed and discussed.

Students will learn 4 techniques:

1. Basic dosimetric measurements and techniques of exposing cells to gamma rays and alpha particles. Students will use low activity gamma radiation sources to measure 1) dose rates in air as a function of distance from the source, 2) the energy spectrum of gamma radiation from ^{137}Cs and ^{133}Ba , 3) the effect of shielding. Dosimetric measurements of high activity ^{137}Cs sources will also be carried out. The nuclide content of a radioactive mineral will be determined by analysing its energy spectrum. Radon activities will be measured with an AlphaGuard meter in air drawn from an encapsulated ^{226}Ra source and in a basement room. Build-up of ^{222}Rn disintegration products will be demonstrated by measuring the increment of gamma radiation in a chamber filled with ^{222}Rn . Techniques of exposing cells to gamma radiation (^{137}Cs) and alpha radiation (^{243}Am) will be demonstrated.
2. Preparation of slides for analysis of chromosomal aberrations as well as microscopic analysis of chromosomal aberrations and micronuclei. A microscope will be available for each student. Slides will contain cells exposed to A) increasing doses of gamma radiation (demonstration of a dose response); B) a same dose of gamma radiation and alpha particles (demonstration of the concept of relative biological effectiveness). Each student will receive slides and images for analysis.
3. In situ hybridisation with whole chromosome probes (FISH) as well as image-based analysis of aberrations in painted chromosomes. Analysis will be done manually on digital images (aim: demonstration of stable and unstable-type aberrations).
4. Detecting gamma H2AX foci and image-based analysis of foci. Analysis will be done on digital images using the Image J software (aim: demonstration of analysis technique taking into account focus size as well as focus distribution).

Experiments will be carried out with CHO cells and human lymphocytes (aberrations and micronuclei) and with RPE-1 cells (gamma H2AX). The cell lines/techniques are established and currently used in our laboratory.

A detailed description of the course is given below. Lectures/demonstrations start at 09:00. Afternoon demonstrations start at 14.00. During week 2, morning exercises start at 09:30.

A detailed description of the course is given below. Lectures will be given by European experts in the field. Lectures will start at 09:00.

Monday - day 1

Morning lecture: DNA damage and repair following irradiation of cells (1.5 h)

Morning lecture: Radiation-induced chromosomal aberrations (1.5 h)

Afternoon: Group 1 learns dosimetry, group 2 learns gammaH2AX detection. Groups 3 and 4 analyse aberrations and micronuclei.

Tuesday – day 2

Morning lecture: AI and its use in radiation research (1.5 h)

Morning lecture: Combined exposures of radiation and other stressors (1.5 h)

Afternoon: Group 3 learns dosimetry, group 4 learns gammaH2AX detection. Groups 1 and 2 analyse aberrations and micronuclei.

Wednesday – day 3

Morning lecture: Factors which influence cellular radiosensitivity (1.5 h)

Morning lecture: Bystander effects of radiation (1.5 h)

Afternoon: Group 2 learns dosimetry, group 1 learns gammaH2AX detection. Groups 3 and 4 analyse aberrations and micronuclei.

Thursday – day 4

Morning lecture: Radiation effects on the immune system and the use of radon to treat autoimmune diseases (1.5 h)

Morning lecture: Statistical analyses of experimental results from low and high throughput approaches in radiation research (1.5 h)

Afternoon: Group 4 learns dosimetry, group 3 learns gammaH2AX detection. Groups 1 and 2 analyse aberrations and micronuclei.

Friday – day 5

Morning lecture: Radiation-induced micronuclei (1.5 h)

Morning lecture: Radiation-induced gammaH2AX foci (1.5 h)

Afternoon: Group 1 learns cell fixation for aberrations, group 2 learns FISH. Groups 3 and 4 analyse aberrations and micronuclei.

Saturday: 09:00 – 17:00 trip to Uppsala with a visit to scientific museums (Lineus, Gustavianum and Karolina Rediviva). 18:00 – 21:00 reception at the university.

Monday – day 6

Morning: (3h) Group 3 learns cell fixation for aberrations, group 4 learns FISH. Other groups analyse experimental results.

Afternoon: All groups learn to score gamma-H2AX foci (4h).

Tuesday – day 7

Morning: (3h) Group 2 learns cell fixation for aberrations, group 1 learns FISH. Other groups analyse experimental results.

Afternoon: All groups learn analyse experimental results (4h).

Wednesday – day 8

Morning: (3h) Group 4 learns cell fixation for aberrations, group 3 learns FISH. Other groups analyse experimental results.

Afternoon: All groups analyse experimental results (4h).

Thursday – day 9

Morning: Collecting and analysing results and preparing presentations – all groups (3h)

Afternoon: Collecting and analysing results and preparing presentations – all groups (5h).

Friday – day 10

Morning: presentation of results, general discussion (3h)

A diagram illustrating the timing of CELET components is shown below. Blue: lectures/seminars, yellow: experimental work in the lab, green: scoring of slides/images.

		Week 1					Week 2				
		Monday	Tuesday	Wednesday	Thursday	Friday	Monday	Tuesday	Wednesday	Thursday	Friday
MORNING	LECTURES	LECTURES					Harvesting Group 3	Harvesting Group 2	Harvesting Group 4	Analysis of results and preparation of presentations	
	LECTURES	LECTURES					FISH Group 4	FISH Group 1	FISH Group 3	Group work presentations	
AFTERNOON	Other groups: aberrations and micronuclei scoring	Other groups: aberrations and micronuclei scoring					Other groups: aberrations and micronuclei scoring	Other groups: aberrations and micronuclei scoring	Other groups: aberrations and micronuclei scoring	Analysis of results and preparation of presentations	
	Dosimetry Group 1	Dosimetry Group 3	Dosimetry Group 2	Dosimetry Group 4	Harvesting Group 1	Scoring gamma H2AX using image J	Scoring aberrations and micronuclei	Scoring aberrations and micronuclei	Free afternoon		
	gH2AX Group 2	gH2AX Group 4	gH2AX Group 1	gH2AX Group 3	FISH Group 2						

	Lectures
	Learning methods in the lab
	Scoring and analysis of results

The course website can be found at: <https://www.crpr-su.se/CELET>