

Single Cell Gel Electrophoresis and Its Applications in Different Fields

Fahim M, Ahmed A* and Hussain S

Department of Pharmacy, Quaid-i-Azam University, Islamabad, Pakistan

Abstract

The Single Cell Gel Electrophoresis assay (SCGE) is a better technique for the detection of DNA damage and DNA repair in eukaryotes. First time comet assay technique was designed by Ostling and Johanson in 1984 and then it was modified by Singh et al. in 1988. There are four main steps in comet assay. These are:

1. Encapsulation of cell in agarose having low melting point.
2. Lysis of cells.
3. Gel electrophoresis and forth is staining and visualization.

This technique is also known as comet assay because DNA movement pattern in gel electrophoresis similar to a comet. The comet assay is use for the measurement of DNA strand breaks in eukaryotes. In this technique cell suspended in agarose on a slide and lysed with detergent to form nucleoid contain supercoiled DNA attached with nuclear matrix. A comet which was made in electrophoresis detected by fluorescence microscopy, manual or mechanical methods use for scoring but now a day's mechanical methods are very useful. Comet assay have simplicity, sensitivity, speed and economical use. In this review we discuss its protocol and its applications in different fields.

Keywords: Gel electrophoresis; DNA; Applications; Comet; Microscopy

Introduction

The comet assay is also known as microgel electrophoresis (MGE) was first time established by Ostling and Johanson in 1984 for the direct exposure of DNA damage in individual cells [1]. First of all cells in a thin agarose gel on a slide (microscopic) were lysed then did electrophoresis and after that stained with a fluorescent DNA binding dye. The electric current pulled the charged DNA from the nucleus such that undisturbed and broken DNA fragments migrated further. The resulting images, which were subsequently named for their appearance as 'comets', were measured to determine the extent of DNA damage [2,3]. The original method which was developed by Ostling and Johanson was not very good because they were unable to remove all of the proteins from the cells. So, more lysis requires for the removal of cellular proteins and for the migration of broken duplex in gel electrophoresis. Two laboratories changed the method about five years ago by applying different denaturing conditions for the measurement of Single strand breaks in DNA [1,3] the first group [3] were maximized the sensitivity for the detection of small number of strand breaks, while second laboratory focused towards the detection of subpopulations which vary in drug or radiation sensitivity [1]. There has been greater interest in comet assay from past two to three years [4-8]. Many papers publish on comet assay because it has very unique design for the direct determination of DNA damage in individual cells. Single cell gel electrophoresis also uses to examine DNA repair by many experimental conditions (Table 1, Figures 1 and 2) [9-15].

Applications

Role of comet assay in genotoxicity testing

Comet assay now a day's used to evaluate the efficacy and safety of pharmaceuticals [16]. It is easily use for *in vivo* experiments; tissues can be spread in a single cell suspension; as well as white blood cells (WBCs) provide the material. More sensitivity as well as extra information we get through comet assay. Insertion of repair endonucleases was used for the measurement of specific lesions. Genotoxicity is also evaluating

in cell culture system, or with the combination of microsomal "S9" taken from liver that provides enzymes to metabolize the chemicals in a more reactive form [10]. The comet assay is famous for the evaluation of phytochemicals ability. For example the protection of cells against genotoxic results [17].

Role of comet assay in chronic and degenerative diseases

Comet assay is used for measure DNA damage in target cells; this DNA damage is very useful for the study of diseases like Diabetes mellitus, Rheumatoid arthritis (a joint disease) and also in the study of Alzheimer's and Parkinson's disease [18]. When we study these diseases through comet assay we seen that high level of SSBs and stress condition were found in a patients, and increased receptiveness as well as efficacy of DNA damage was decreased [19]. The DNA damage in diseases like diabetes, Alzheimer's etc. is due to increased oxidative stress or due to direct effect on DNA by different toxic chemical compounds. Alkaline comet assay combined with measurement levels of 8-hydroxyl-2-deoxyguanosine was very suitable method for the evaluation of DNA damage due to slight oxidation.

Use of comet assay in male infertility test

Now a day's more emphasis has been shift towards sperm DNA damage which causes infertility in males. When we study quality of sperm DNA damage in patients we seen that more sperm DNA damage occur in patients having infertility as compares to fertile men, the cause of sperm DNA damage is due to endogenously derived reactive oxygen

***Corresponding author:** Ahmed A, PhD Scholar, Department of Pharmacy, Quaid-i-Azam University, Islamabad, Pakistan, Tel: 923338619923; E-mail: aliahmed@bs.qau.edu.pk

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S No.	Procedures	Advantages	Disadvantages/limitations
1	PCR	Easy to measure gene specific DNA damage	Cannot quantify
2	TUNNEL	Detect single and double strand brake	Cannot differentiate between necrosis and apoptosis
3	HPLC	Sensitive and specific for quantification of thymidine	Early elution property of liquid chromatography
4	Micro-nucleus assay	Easy to perform less time required	Low sensitivity
5	Halo assay	Detect changes in DNA of individual cell	Low sensitivity
6	FISH	More sensitive than FCM	well-resourced laboratory facility requires
7	FCM	Detect DNA damage in apoptotic cells	Low sensitivity as compared to FISH
8	GC-MS	Sensitive to detect DNA damage	more estimation of damage
9	Immunological assay	Low amount of DNA requires	Costly equipment
10	Comet assay	Detect DNA damage in individual apoptotic cells as well as differentiate between apoptosis and necrosis	Small deletions cannot be detected

Table 1: Comparison between comet assay and different techniques.

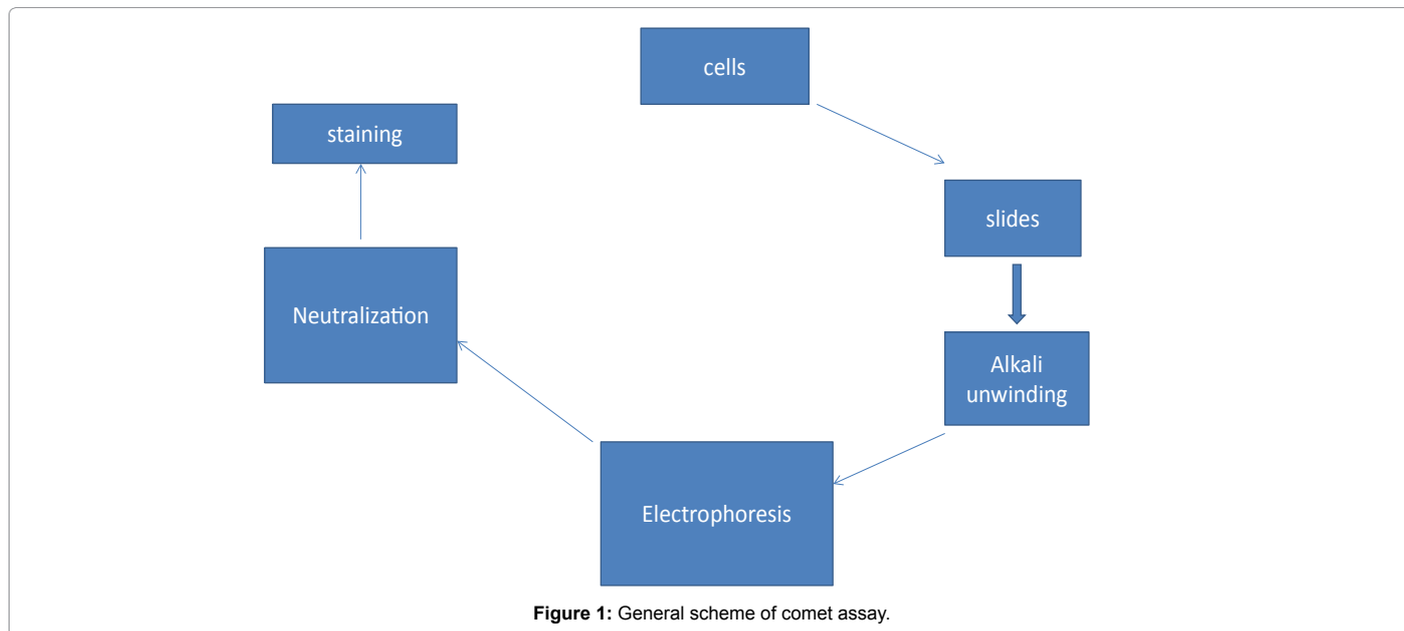


Figure 1: General scheme of comet assay.

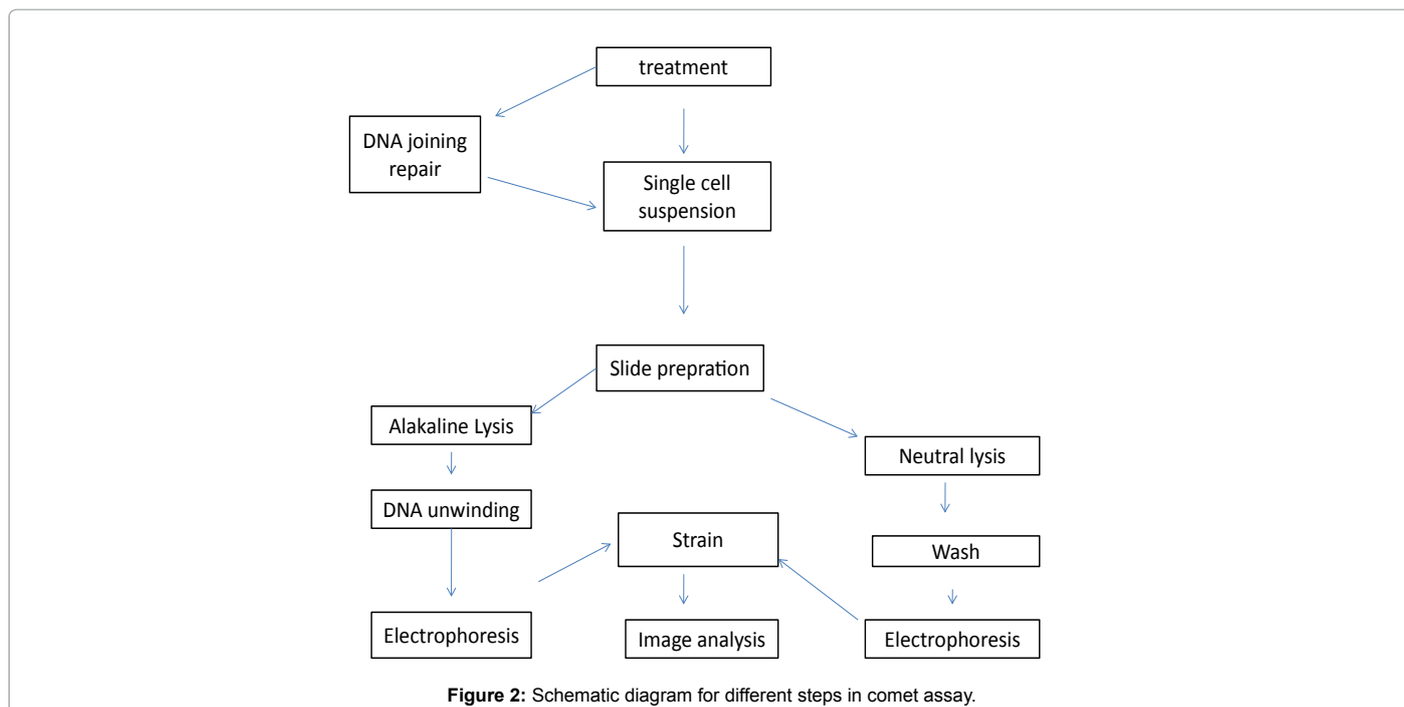


Figure 2: Schematic diagram for different steps in comet assay.

species from polyunsaturated fatty acids [20]. Comet assay was found very sensitive technique in the evaluation of DNA damage as compare to other techniques like TUNNEL, HALO assay, flow cytometry etc. [21]. We mention Table 1 in which comparison of comet assay with different techniques has been discussed. The challenges which we are facing while using comet assay for the assessment of sperm DNA damage, are that DNA have a unique structure as compare to other cells. So we need to modify comet assay technique and optimizing conditions.

Further study now a day for understanding the mechanism of paternal DNA is under consideration and scientists working on it [22].

Comet assay role in ecogenotoxicology

Ecogenotoxicology is basically the study of environmental effects by genotoxin. Genotoxin chemicals can damage genetic material (DNA) of humans and also affects other organisms. Genotoxin chemicals cause pollution in an environment and badly effect the environment. There is a difference between the study of environmental toxicology and epidemiology former ones has focus on species or population level but the later focus on individuals. Some organisms selected as indicators for the detection of toxins for example radioactive compounds, heavy metals which cause harmful effects on human population. The organisms which are selected as indicators are snails, mussels, fish, mice, and some plants as well. Comet assay is use for the monitoring of toxic effects of above mention species [23]. There are many ways for investigation like to carry out laboratory experiments, organism expose by given them DNA damage agents for the study of sensitivity shown by selected organisms. Specimens should be placed in a clean location so that analysis of DNA damage would be easy.

Comet assay role in genotoxicity of nanoparticles

Nanoparticles in the past were used in research, development and in commercialization, but they had toxicological issue they damage DNA. Lots of mechanisms involve in the ability of nanoparticles for the damage of DNA. These nanoparticles cause oxidative stress but some mechanisms tells us that direct Nanoparticles and DNA connections occur, and causes trouble in mitotic spindle and its components Size, discharge of toxic ions and ability to generate ROS are the physicochemical properties important for DNA damaging effects [24]. Nanoparticles can also cause genomic instability by epigenetic mechanisms. There are many assays available which can be used to detect Genotoxicity of nanoparticles. Comet assay help in the measurement of DNA damage as well as DNA repair.

Comet assay use in the study of cancer

Comet assay is very helpful in the evaluation of carcinogens/chemicals. *In vivo* comet assay is use to differentiate between genotoxic and non genotoxic chemicals which are used in the field of genotoxicology. Comet assay use biomarker and detect DNA damage which is the leading cause of carcinogenicity. Comet assay is a better method as compare to other techniques use for assessment. DNA damage and defect in DNA repair mechanism cause initiation of pathogenesis and development of cancer. The patients which have cancer, in them increase level of DNA damage and defective repair mechanism can be seen [19]. By finding the level of DNA damage we get information about the nature and harshness of cancer which is helpful for the treatment of cancer and for curing patients from cancer.

Forecast of tumor radio and chemo sensitivity

Treatment of cancer depends upon the nature of tumor cells in a

patient body. In the past for the study of tumor cells sensitivity towards radiation clonogenic cell survival assay, image of tumor cell by PET scan and single photon emission CT were used. But above mentioned techniques require more time. So we use Comet assay for measuring the sensitivity of tumor cells towards radiation and also in a cells having low proliferation. Alkaline comet assay could be very helpful for the study of different tumor cell lines like Colon breast prostate. Alkaline comet assay is also very helpful in the prediction of radio sensitivity of tumor cells and also determine their efficacy. Comet assay is also a sensitive technique for evaluating the use of chemotherapeutic agents in the treatment of cancer [15,25].

Diagnosis

Comet assay can also be used for diagnosis purpose. For example in a disease named as Nijmegen breakage syndrome which is an autosomal recessive disorder in which genetic instability and cancer occur. With the help of comet assay it is easy to identify Nijmegen syndrome because high level of strand breakage in lymphocytes. Second disease which is identify through comet assay name as xeroderma pigmentosa. In this disease defect occur in nucleotide excision repair system. Normal UV induced accumulation of DNA strand breaks does not seen in xeroderma pigmentosa, so comet assay use [26].

Limitations of Comet Assay

Comet assay is a sensitive and better technique for DNA damage and also in measure of DNA repair but comet assay also have some limitations. 1. Small number of sample can be run in a single electrophoresis; it can hold only 20 slides and every slide contain one or two gels. Now changes have been made in comet assay by reducing the gel size. Automated scoring now requires as compare to simple manual scoring [27].

Conclusion

Single cell gel electrophoresis is a simple and rapid method for the study of DNA damage and DNA breaks, now plenty of modified versions available and wide range of cells arranged in the comet assay, makes it very helpful tool for the estimation of extent of DNA damage and repair in various diseases like cancer, Alzheimer, Parkinson's etc. and also in male infertility. It is very sensitive and better technique as compares to other technique like TUNNEL, HPLC etc.

Conflict of Interest

We declare no conflict of interest.

Source of Funding

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