Fluorometric Quantitation of dsDNA Using PicoGreen®

Abstract

An essential element of cellular and molecular biology is the ability to quantitate DNA in large numbers of samples at a sensitivity that enables determination of small amounts of sample. Here we describe a method to quantitate dsDNA using the BioTek Synergy™ HT and Synergy™ 2 Multi-Detection Microplate Readers.

Introduction

Many techniques in cellular and molecular biology require the ability to quantitate dsDNA in large numbers of samples at sensitivities that only require a small amount of the total sample. Isolation of plasmids from bacterial cultures, genomic DNA from mammalian cells, cDNA synthesis for library production, and quantitation of PCR products for diagnostic purposes all require the direct quantitation of dsDNA. Also, many biochemical studies that involve the growth kinetics of cell cultures or cell cycle studies require normalization by DNA content.

The most commonly used method for the determination of nucleic acid concentration is the determination of absorbance at 260 nm ($A_{260}$) as described by Matiatis et al [1]. This method, while quite adequate for many situations, can suffer from the interfering absorbance of contaminating molecules. Many of these contaminants, which include nucleotides, RNA, EDTA and phenol are commonly found in nucleic acid preparations. As a result, several fluorescent staining techniques have been developed to alleviate many of the problems associated with absorbance at 260 nm [2-4]. One such stain is PicoGreen®, which in conjunction with the BioTek Synergy™ HT and Synergy™ 2 Multi-Detection Microplate Readers offers high specificity, as well as, high sensitivity for dsDNA quantitation.

Materials and Methods

A PicoGreen dsDNA Quantitation Kit, catalogue number P-7589, was purchased from Invitrogen (Carlsbad, California). The solid 96-well black microplates, catalogue number 3915, were purchased from Corning (Cambridge, Massachusetts).

A series of dilutions ranging from 0.0 to 1000 ng/ml of purified Lambda DNA was made using TE (10 mM Tris, 1 mM EDTA, pH 7.5) as the diluent. Working PicoGreen reagent was prepared by diluting the concentrated DMSO-PicoGreen stock solution, provided in the PicoGreen kit, 1:200 with TE according to the kit instructions (2). Equal amounts of DNA sample and working PicoGreen dsDNA quantitation reagent were mixed and incubated for 5 minutes at room temperature. Following incubation, 200 μl aliquots were pipetted into microplate wells in replicates of eight. Fluorescence was determined using either a BioTek Synergy HT or a Synergy 2 Multi-Detection Microplate Reader. Both readers used a 485 nm, 20 nm bandwidth.
excitation filter and a 528 nm, 25 nm bandwidth emission filter. In addition, the Synergy 2 used a 510 nm cutoff dichroic mirror. The reader was programmed to autoscale the sensitivity setting to the highest DNA concentration. The data was collected from the top using a 10 msec delay after plate movement, 50 reads per well. Although in these experiments the plates were read immediately, if they remained sealed and protected from light the reaction was found to be stable for several hours.

Results

Fluorescence intensity was determined for DNA concentrations ranging from 0 to 1000 ng/ml. Over this range the intensity increased in a linear fashion. Using Gen5™ Data Analysis Software (BioTek Instruments), a least means squared linear regression analysis can be generated with a coefficient of determination ($r^2$) value of 1.0. The average coefficient of variance (%CV) of the standards was less than 3%, with the greatest variation in the lower DNA concentrations tested (data not shown). In terms of sensitivity, the assay was found to be sensitive to the picogram level. Under appropriate sensitivity settings, DNA concentrations as low as 50 pg/ml were found to be statistically different (P<0.05) from the TE only, 0 ng/ml, control. Quantitation of dsDNA using the fluorescent properties of PicoGreen in conjunction with the BioTek Instruments’ readers allows researchers to quantitate as little as 10 pg/well (50 pg/ml in a 0.2 ml total volume) thus providing reliable quantitation of dsDNA concentrations ranging over four orders of magnitude.

![Lambda DNA Concentration Curve](image)

**Figure 1. Lambda DNA Concentration Curve.** The fluorescence of Lambda dsDNA concentrations from 0 to 1000 ng/ml stained with PicoGreen was measured using a Synergy™ 2 Multi-Detection Microplate Reader. Linear regression analysis was performed using Gen5™ Data Analysis Software.
Figure 2. Lambda Low DNA Concentration Curve. The fluorescence of Lambda dsDNA concentrations from 0 to 625 pg/ml stained with PicoGreen was measured using a Synergy™ HT Multi-Detection Microplate Reader. Linear regression analysis was performed using Gen5™ Data Analysis Software.

Very similar results were obtained with both the Synergy HT and the Synergy 2 Multi-Detection Microplate Readers. Due to the use of dichroic mirrors in addition to specific wavelength bandpass filters the Synergy 2 reader is typically more sensitive than the Synergy HT reader. The dichroic mirrors reduce light cross talk between the excitation and emission filters that can occur with filter pairs that are close in wavelength. The net effect of this is to reduce background signal, which results in lower detection limits. The fact that both readers, the Synergy 2 with dichroic mirrors and the Synergy HT without, have virtually the same detection limit suggests that the detection limit we are observing is that of the chemistry rather than the reader.

In order to achieve greater sensitivity, measures can be undertaken to reduce background fluorescence. Removal of certain common contaminates can lead to an increase in sensitivity. Several compounds decrease the fluorescent signal when present in a PicoGreen-DNA assay. In particular, the presence of 200 nM sodium chloride or 50 mM magnesium chloride can decrease the signal by approximately 1/3. In cases where only low DNA concentrations are to be determined, a calibration curve utilizing lower concentrations can be used in conjunction with setting the fluorescence reader’s gain such that the highest DNA concentration yields a fluorescence intensity near the instrument’s maximum. Alternatively, the use of different microplates that utilize lower fluorescent plastics would be expected to reduce background and therefore increase sensitivity.

The ability to perform this assay in microplates offers several advantages over the conventional tube-based fluorescence assays. Like most assays that are performed in microplates, the ability to use multi-channel pipettes greatly reduces the manual labor required to perform the assay. The microplate format also lends itself to “off the shelf” automation for laboratories with high volume requirements. The smaller reaction volumes in microplates will lead to lower per assay costs by reducing the amount of expensive reagents necessary to perform the assay.
References


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*Rev. 12/1/06*