

Compared between sensevityof bacteria streptococcus spp. For some antibiotic by using semi logarithmic plots

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Abstract—In this paper three sorts of anti-toxin for information sensevity of microscopic organisms streptococcus spp. Confined from patients contaminated by tonsillitis, and looked at between microscopic organisms sensevity for this sorts of anti-toxin to three arrangement contain every one of it ten million from microbes cells on various period time. We were taking when deciphering the coefficients of sham variables in semi-logarithmic relapse models. Existing results in the writing give the best-impartial estimator of the rate change in the needy variable, inferred by the coefficient of a fake variable, and of the fluctuation of this estimator. We develop these outcomes by setting up the definite testing appropriation of an unprejudiced **estimator** of the suggested rate change. This appropriation is no ordinary, and emphatically skewed in little specimens.

Key Words—semi logarithmic plots; Semi-Log Model; Semi-Logarithmic Graphs; Logarithmic function; Exponential growth; Detection time; logarithmic order of death.

INTRODUCTION

Science works by building 'models'. Not minimal cardboard and plasticine models, but rather models made out of images. We can play with the typical models and modify them until they begin to carry on as it were, which looks like the things we think about. When we have done this, we get a comprehension of the things we think about which is much more profound than we would ever get on the off chance that we adhered to words and pictures. Scientific models don't supplant words and pictures, they hone them, So models extend our comprehension of 'frameworks', whether we discussing a system, a robot, a synthetic plant, an economy, an infection, an environment, a growth or a mind. Moreover, important to comprehend something about how models are made. There is a

colossal scope of valuable models attacking the Life Sciences[Richard D.1976],[Richard D. 1986] , [Richard D.1989].

In the 80's, with effective desktop processing getting to be regular utilize, the name 'prescient microbiology' was instituted for this region of numerical microbiology. Its nearest proportional can be found in biotechnology with its numerical models very much expounded in the 60's. At the point when the author of this theory [JózsefBaranyi 2010], began his work in this field, thoughts in regards to microbial development models connected in biotechnology were taken from the great papers of Tsuchiya et al (1966), Frederickson et al (1967), Turner et al (1976), Roels and Kossen (1978), Srivastava and Volesky (1990), Nilsen and Villadsen (1992).

Microscopic organisms matter rather a great deal, since some of them live on, or in, us. Of around fifteen hundred known types of microscopic organisms, somewhere in the range of two hundred can make you be wiped out. Just a century prior, making due to age five was a chancy business, and it may turn out to be so once more. Regardless it is in numerous parts of the world. So understanding the things is of some squeezing significance. Models for comprehension the development of microscopic organisms are likewise valuable in comprehension the development of different things, for example, your bank equalization or little organizations. So kindly don't switch your mind off on the grounds that you are not a scientist [Michael D. 2001].

The instructional exercise on the relations between populace information and the rates of development that computed from information where semi-logarithmic diagrams will be talked about as a gadget for speaking to the extent of developing populaces and for breaking down the way of the development [Albert A. 1993].

An essential methodology in microbiology is the investigation of development tentatively normally basic since development (engendering) of microorganisms can be quick (in contrast with higher creatures) and requires just little culture volumes (jars) [Friedrich W. 2010]. A few creators and analysts say; [Kiryl T., 2010] contemplated exponential development and rot in common marvels, (for example, populace development, radioactive rot, and so forth.) amounts develop or rot at a rate corresponding to their size, [David E. 2011] concentrated on translating sham variables in semi-logarithmic relapse models: careful conveyance comes about; he broaden these come about by building up the definite testing appropriation of a fair-minded estimator of the inferred change. He talk about the development of bootstrap certainty interims for the inferred rate change (ewp). A general hypothetical structure to examine the shape elements of effectively developing and redesigning surfaces. Utilizing system to build up a physical model for developing bacterial cell dividers and study the interchange of cell shape with the progression of development and choking, foresee that exponential development in cell size requires a steady measure of cell divider vitality to be disperse per unit volume [Shiladitya B. and et al., 2016].

The one of arise in many areas of science that the Logarithmic graph the desirability of working directly with measuring comparable that is specialty with studying the comparison between sizes of the different types for beings alive. By supposed that the two variables x, y size of two different parts from the bodies of individuals of beings alive, for example; x will represent the average weight for the body. When y represent the average weight, grey matter for individuals have the different age belongs to human gender, whenever individual change in age; the average amount change for x and y. and they often found associated by relation on the form $y = ax^b$ so it's called power function and these function known by Huxley's Law for simple measuring comparable. This law usually used to compare measure between different types of beings alive. This is what has been discuss in this research.

For experiment, the set of measures flawed simple measuring comparable that must be used the Logarithmic graph for variable x versus variable y.

Finally, the result are either given in tabulated form or illustrated in figures, Adoption of MATLAB program in order to give a good comparison for result and also to be more interest to the reader.

Semi-Log Model [David M., 1999]:

The slop coefficient measures the relative change in Y for given supreme change in the estimation of the logical variable (t).

By using calculus:

$$\begin{aligned}
 b_2 &= \frac{\partial \ln Y}{\partial t} \\
 &= \left(\frac{1}{Y}\right) \left(\frac{\partial Y}{\partial t}\right) \\
 &= \frac{\frac{\partial Y}{Y}}{\partial t} \\
 &= \frac{\text{proportional alteration in } Y}{\text{proportional alteration in } t}
 \end{aligned}$$

In the event that we increase the proportional alteration in Y by 100, we get the rate or development rate in Y for a flat out change in t.

The slope coefficient measures the instantaneous rate of growth

r is the compound growth rate

Where $b_2 = \ln(1 + r)$

Antilog (b_2) = $(1 + r)$

So $r = \text{antilog}(b_2) - 1$

So $r = \text{antilog}(0.0269) - 1$

$r = 1.0273 - 1 = 0.0273$

If $b_2 > 0$, then an upward direction. Where if $b_2 < 0$, then a downhill direction

How microbes die: the logarithmic order of death [SHTM 2010 (pt.5) Sterilization]:

Life forms which kick the bucket as a consequence of a forced anxiety bite the dust in a systematic, and unsurprising, way. This can be spoken to as survivor, demonstrating the quantity of living beings as yet living at different times after the start of introduction to the anxiety condition.

Whichever multicellular life forms are tried, for instance creepy crawlies or plants, and whatever the deadly stretch, the survivor bend remains basically the same. This was acknowledged as generally valid for all living beings until the mid 1900s when laborers, for example, Harriet Chick [see Chick (1908)] demonstrated that in a homogeneous society of a solitary strain of microscopic organisms the phones kicked the bucket at a steady rate when presented to a specific deadly push.

Then again the quantity of survivors might be plotted on a logarithmic scale as a component of time on the number-crunching scale, which is

alluded to as a semi-log survivor bends (see Fig.(1) and (2)). While both the number-crunching and semi-log survivor bends precisely speak to the passing of microscopic organisms the last is more valuable in sanitization considers where interest is focused on the rate of pulverization as the quantity of survivors methodologies zero.

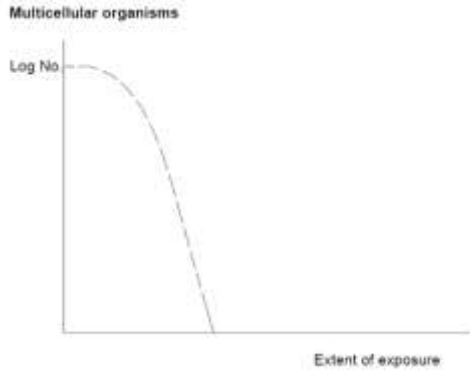


Fig (1) Semi-log survivor curve for multicellular organisms

It is normal to utilize the last approach since in sanitization contemplates we are keen on the rate of decimation as the quantity of surviving small scale living beings approaches zero, which is best demonstrated utilizing a logarithmic plot.

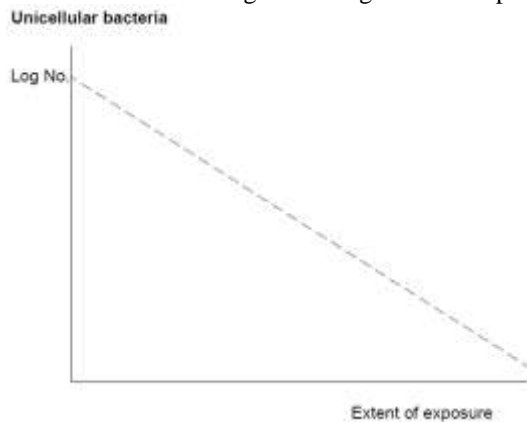


Fig (2) Semi-log survivor curve for unicellular bacteria

Experience has demonstrated that the semi-log survivor bend for warmth push frequently approximates to a straight line for part or the greater part of the survivor bend. However there are numerous recorded cases where deviations from the "perfect" straight line condition happen.

Treatment of sterilization-process microbial survival data[SHTM 2010(pt.5) Sterilization]:

A mathematical way to deal with the resistance of microscopic organisms to warm passing is require to permit estimation of identical lethality. Two elements should be viewed as; the

warm resistance of the miniaturized scale life form at a specific temperature and the adjustment in that resistance which happens with changes in temperature.

These two variables are comparable to the rate consistent and temperature coefficient of a substance response, individually.

Spore inactivation in sodden warmth might be considered as a monomolecular first request response, that is the place the rate of response is administered by the centralization of the reactant, for this situation the bacterial spores.This may be expressed as

$$\frac{dN_a}{dt} = kC_a$$

Where t = time

Ca = spore concentration

k =a reaction rate constant at constant temperature

$$\text{Then } (\log C_a^0 - \log C_a) = k(t - t^0)$$

Where the superscript 0 indicates initial conditions.

A semi-logarithmic plot of fixation versus time will yield a straight line of incline k. k has measurements of time-1. The negative proportional of the rate consistent k is equal to the quantity of minutes required to inactivate 90% of the life forms exhibit, that is a 1 log lessening. This worth is alluded to as the D esteem and, as expressed, numerically it is contrarily corresponding to the inactivation rate steady k.

$$D=2.303/k.$$

Semi-Logarithmic Graphs [Albert A. 1993]:

On a semi-logarithmic chart, a straight line speaks to enduring development. This is critical in perceiving regardless of whether a progression of information focuses speak to relentless development. To see this property, let us take the normal logarithms of both sides of eq.

$$\ln N = (\ln N_0) + kt$$

If we plot (ln N) vs. t we have a straight line whose slope is k and whose intercept at time t=0 is (ln N₀).

One uses an exploratory adding machine to "gaze upward" the estimations of the logarithm. Common logarithms are rung utilizing the "ln" key,

and logarithms to the base ten are rung with the "log" key. Either kind of logarithm can be utilized.

Semi-Logarithmic Graphs Paper [Albert A. 1993]:

The most advantageous approach to make a semi-log plot is to utilize semi-log diagram paper. This imprinted in a manner that separations on the vertical scale are corresponding to the logarithms of the amount being plotted, while the level scale is the typical direct scale. This spares one the need to turn upward logarithms.

Critical properties of semi-log paper are that the greater part of the many years of progress on the vertical scale are spoken to by the same separation, and there is no zero on the vertical scale. In this manner, the separations on the vertical scale are the same for the numbers from 1 to 10, as from 10 to 100, as from 100 to 1000, and so on. Semi-log chart paper, one must determine what number of cycles one needs. "One-cycle" paper will oblige one decade of information; "two-cycle" paper will handle two many years of information, as from 1000 to 100,000. "Five-cycle" paper will suit information extending more than five decades.

Exponential growth for many doubling times [Albert A.1993]:

An imperative component of consistent development is that after drawn out stretches of time (numerous multiplying times), the span of the developing amount gets to be huge.

For mental figurings, it is helpful to recollect that

$$2^{10} = 1024 \approx 10^3$$

So the growth in 25 doubling times can be estimated as follows:

$$2^{25} = 2^{10} \times 2^{10} \times 2^5 \approx 10^3 \times 10^3 \times 32 = 3.2 \times 10^7$$

Detectiontime [József B 2010]:

Practically speaking, it is hard to take after the division of individual cells with computerized estimations. Assume that $y_{det} = \ln(x_{det})$ is the level at which a homogeneous bacterial society can be recognized, for instance, by measuring its turbidity in a fluid medium. Assume that y_{det} is in the exponential stage and let T_{det} mean the identification time at which the single-cell produced subpopulation achieves this y_{det} limit esteem. At that point the identification time is a moved adaptation of the L_g parameter. Be that as it

may, this quality relies on upon the quantity of starting cells and in the event that this is obscure, it is difficult to surmise to the single cell slack times.

Our fundamental goal is to utilize discovery time information to give an exact estimation technique for the appropriation parameters of the single cell slack times, when the underlying cell number is arbitrary. We will likewise examine some functional ramifications with respect to Quantitative Microbial Risk Assessment.

Bacterial survival curves [József B 2010]:

We have visible that for nourishment security demonstrating, it is indispensable to study and model the adjustment time when microscopic organisms are immunized in another environment. The circumstance is comparative with survival bends where a purported "shoulder" period goes before the exponential rot. Mirror pictures of development capacities are likewise used to model survival bends (Xiong et al., 1999b) yet a parallel study on development and survival models delineated in this part demonstrates the symmetry is not that clear as it takes a gander at the main sight.

For both development and existence models, we recall the "ln tallies v. time" bend the populace bend. At the point when the phones' surroundings all of a sudden gets to be inactivating, (for example, warm therapy of sustenance), the "opportunity to death" for a solitary cell will be known as the individual survival time.

The primary inquiries to be replied in this part are: What is the relationship between the appropriation of individual survival times and the survival bend of the populace; besides between the circulation of individual slack times and the development bend of the populace.

We are especially intrigued by creating formulae between the parameters of the circulation elements of individual survival/slack times and the parameters of the survival/development bends of the populace. For this reason, we present the accompanying definition. On the off chance that the development/survival bend (implied as the "ln checks v. time" bend, $y(t)$) asymptotically unites to a straight capacity $y_a(t) = \ln N + \mu(t - \lambda)$

where N is the initial counts, μ and λ are constant and do not depend on N ,

(i.e. $|y(t) - y_a(t)| \rightarrow 0$ as $N \rightarrow \infty, t \rightarrow \infty$), then μ and λ are called the limit slope and limit shoulder / lag parameter of the survival/growth curve, respectively (see Fig.3).

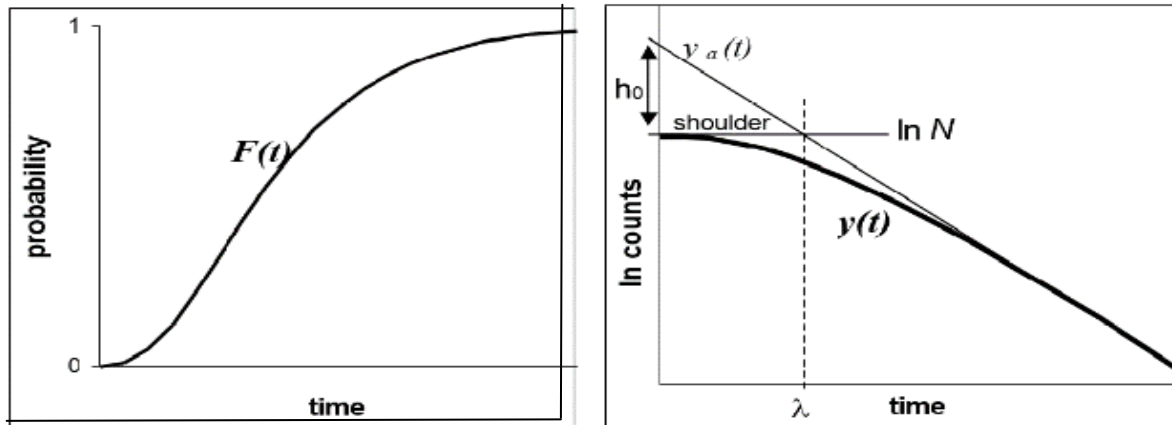


Fig.(3). There is a coordinated charting between $F(t)$, the aggregate dissemination capacity of the survival times of individual cells and $y(t)$, the survival bend of the populace. On the off chance that $y(t)$ asymptotically joins to a direct capacity, $v_{\alpha}(t)$, then a shoulder parameter, λ , is translated for the populace bend. The parameter h_0 can be translated as in the development situation

$$\frac{dq(t)}{dt} = \mu(E(t))$$

Indicate the survival time of the i -th cell, amid an inactivation procedure, by τ_i ($i=1 \dots N$). As some time recently, for similarity with the separate distribution (Baranyi and Pin, 2001), we compare τ_i with the "geometrical" meaning of slack, characterized by a bi-phasic capacity fitted to the logarithm of the produced subpopulation.

Simulating bacterial growth under dynamic conditions [József B 2010]:

A standout amongst the most critical elements of the created model is that the slack is considered as an alteration procedure, where the rate of change is v_b . On the off chance that the E genuine environment changes amid the slack time, while still stays in a development supporting district, then it is a sensible presumption that the $v_b(E)$ and $\mu(E)$ quickly take up the qualities normal for the real environment. The "safe place" of this suspicion was examined by Baranyi et al (1995).

The presence of mathematical arrangement was noteworthy for information fitting. In any case, if nature changes amid the bacterial reaction, mathematical arrangement for the most part does not exist. With the supposition above, we can in any case utilize the first ODE structure (for effortlessness, with the $v_b = \mu$ presumption):

$$\frac{dx(t)}{dt} = \frac{q(t)}{1 + q(t)} \mu(E(t)) x \cdot u(x)$$

Baranyi et al (1995) tried this model. Singular development bends produced under various yet consistent E situations were fitted by the logarithmic structure, from which a $\mu = \mu(E)$ alleged auxiliary model was set up. As have been specified, $h_0 = \mu_{max} \cdot \lambda$ could be taken as a consistent, from which the q_0 beginning quality was figured. The conveying limit of the earth was likewise taken as steady (this amount does not have criticalness in nourishment microbiology at any rate, in light of the fact that when microscopic organisms achieve the conveying limit of the earth, the sustenance is unpalatable at any rate).

The outcome is shown in Fig.(2). The earth was described by temperature as it were. The reenactment anticipated the bacterial development exceptionally well, notwithstanding amid the conformity (slack) period. This forecast would not have been conceivable with previous prescient models utilized as a part of prescient microbiology (see a rundown of those in Zwietering et al, 1990). The key was the dynamic (ODE) foundation and the presentation of the physiological state described by a basic substance that is bit by bit enhancing at the same rate as the particular rate.

Note that the $\alpha_0 = \text{const}$ supposition turned out to be substantial just in a "smooth" development supporting locale of the temperature (see the first distribution). What's more, late studies have demonstrated that the presumption does not hold for other natural elements, similar to pH.

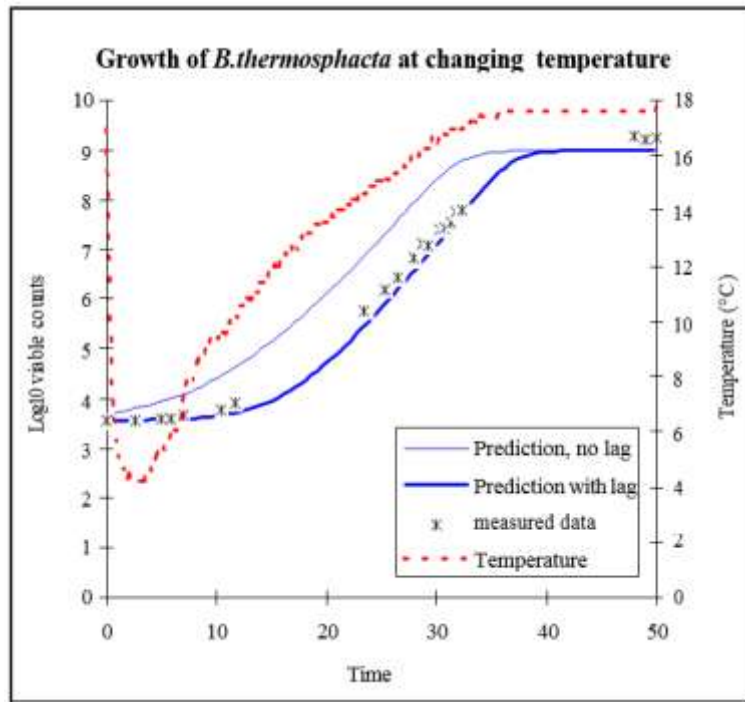


Fig. (4). Bacterial development when the temperature changes (amid the slack time, as well). The temperature profile (spotted red line) is recorded in a cooler, the entryway of which was left open after 2hrs, so the nourishment put inside initially chilled off then expanded back to surrounding. The information (dark stars) are not fitted but rather anticipated (thick blue line) from parameters measured at consistent temperatures. For exhibit, the flimsy blue line is the anticipated bacterial development, had the physiological state been 100% reasonable to the new environment (i.e. $\alpha_0=1$; $h_0=0$: the no-slack circumstance.)

Semi-logarithmic Plots [Jagdich C. and Robin W. 1979]

Semi logarithmic plots used used when the variables associated form exponential function

$$y = ba^x \dots\dots\dots(1)$$

There is another kind of plots for used when the variable y is an exponential function with respected to x, the two variables are associated by using the form of equation $y = ba^x$.

Where a, b are steady requirable to agreeing from the arrangement of trial information, once coordinate representation for a variable x versus variable y prompted getting an a bend. At the point when $a>b1$ then this bend speak to exponential development or expanding development while if $a<1$ that is directed to exponential rot. Extra steady b it has essentially spoken to coefficient change the estimation bend sketch in bearing capitated. By taking Logarithm of both sides of equation (1) one can get:

$$\log y = \log (ba^x) = \log b + \log a^x = \log b + x \log a \dots\dots(2)$$

By substitute $Y=\log y$ one can have

$$y = (\log a)x + \log b \dots\dots\dots(3)$$

Therefor if we sketch Y versus X that is make the position of experimentalism points on or near to the straight line. The slope of straight line is $\log a$ the abscissa y to the intersection point with y axis represented by $\log b$ therefore by measure the slop and spotting the abscissa to the intersection point one can specifying the value of each constants a, b ; the graph for variable $Y = \log y$ versus x named semi-logarithmic Plots.

Sources of data collection :

Applied this model to patients with recurrent tonsillitis caused by bacteria and gram positive as study included 103 patients reviewed the Rizgari Teaching Hospital in Erbil for the purpose of tonsillectomy.

Source: thesis is titled "immunological study of patients with rheumatoid recurrent bacterial tonsillitis" provided by "Taban K. Rashid 2007" submitted to Council of the College of Science - University of Babylon, In Partial Fulfillment of the Requirements for the degree of Doctor of Philosophy in life sciences-biology microscopic requirements.

Analysis and testing of models and discuses results:

By adding bacterial antibiotic to a solution contain each one it ten million from bacterial cells on different period time and then calculating the numbers of bacterial cells that kept life and by continuing in this manner one can get the tabulated results. The following table gives the comparison

between the results that obtained from adding three bacterial antibiotic to a same type of bacteria cells on different period time.

table (1): the number of cells before and after adding the bacterial antibiotic

i	Time in minutes t_i	the number of cells before $x(t_i)$	the number of cells after adding the first bacterial antibiotic $f(t_i)$	the number of cells after adding the second bacterial antibiotic $w(t_i)$	the number of cells after adding the third bacterial antibiotic $z(t_i)$
1	0	10^7	10^7	10^9	10^5
2	10	3.2×10^6	3.2×10^6	3.2×10^8	3.2×10^4
3	20	10^6	10^6	10^8	10^4
4	30	3.2×10^5	3.2×10^5	3.2×10^7	3.2×10^3
5	40	10^5	10^5	10^7	10^3
6	50	3.2×10^4	3.2×10^4	3.2×10^6	3.2×10^2
7	60	10^4	10^4	10^6	10^2
8	70	3.2×10^3	3.2×10^3	3.2×10^5	3.2×10
9	80	10^3	10^3	10^5	10

By using, the semi-logarithmic Plots, one can find the relation between t_i and $f(t_i)$ that represent adding the first antibiotic, relation between t_i and $w(t_i)$ that represent adding the second antibiotic, relation

between t_i and $z(t_i)$ that represent adding the third antibiotic.

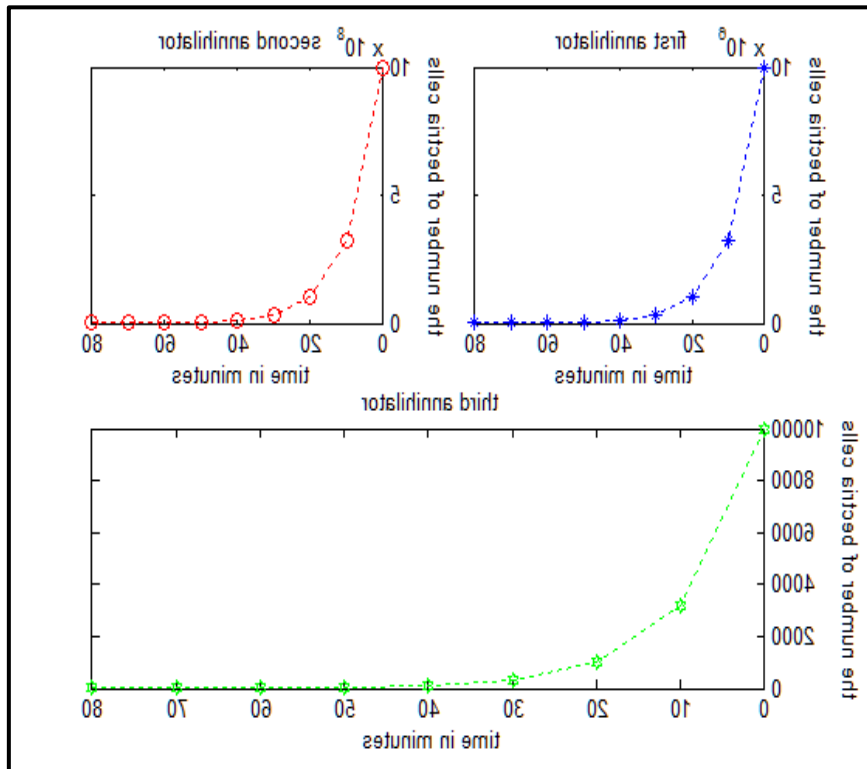


Fig. (5) the relation between the time and number of bacterial cells when adding the antibiotic bacterial.

When sketch $f(t_i)$, $w(t_i)$, $z(t_i)$ versus t the points of experimental data explanation of exponential decay from the type $y = ba^x$ (where $a < 1$) that is may be

sufficient to suitable the data and in this case we will use the semi-logarithmic Plots and finding the two constants a and b .

table (2) taking the logarithm to number of bacterial cells after adding the antibiotic bacteria

<i>i</i>	Time in minutes t_i	the logarithm to number of bacterial cells after adding first antibiotic bacteria	the logarithm to number of bacterial cells after adding second antibiotic bacteria	the logarithm to number of bacterial cells after adding third antibiotic bacteria
1	0	16.1181	20.7233	9.2103
2	10	14.9787	19.5838	8.0709
3	20	13.8155	18.4207	6.9078
4	30	12.6761	17.2812	5.7683
5	40	11.5129	16.1181	4.6052
6	50	10.3735	14.9787	3.4657
7	60	9.2103	13.8155	2.3026
8	70	8.0709	12.6761	1.1632
9	80	6.9078	11.5129	0

When we assign the value of $Y = \log y$ that we get it from the real data one can notice that the point's analogue that denoted by squares with respected to the first antibiotic and denoted by stars with respected to the second antibiotic and asterisk with respected to the third antibiotic. Then the slope of straight line for each cases gives (0.1151) and this will equal to $\log a$.

That is mean $a=0.1220$ when the y abscissa for intercepts point with y-axis is 16.1181 and represent by $\text{Log} b_1$ realized that $b_1=1.000e+007$ and one can get the relation $y=1.000e+007(0.1220)^t$ represent the change in number of cells after adding the first antibiotic.

Another y abscissa for intercepts point with y-axis will represent by $\log b_2=9.2103$ so that $b_2=9.210$ and one can have the relation $y=9.210e(0.1220)^t$ which represent the change in the number of cells after adding the third antibiotic.

There was another method for sketch the semi-logarithmic plots. Not necessarily to compute the value of $\log y$ because instead of that one can used a special kind of paper named as a paper of graph semi-logarithmic Plots which just one of gradation will become logarithmic and the other assign to simple ordinary gradation.

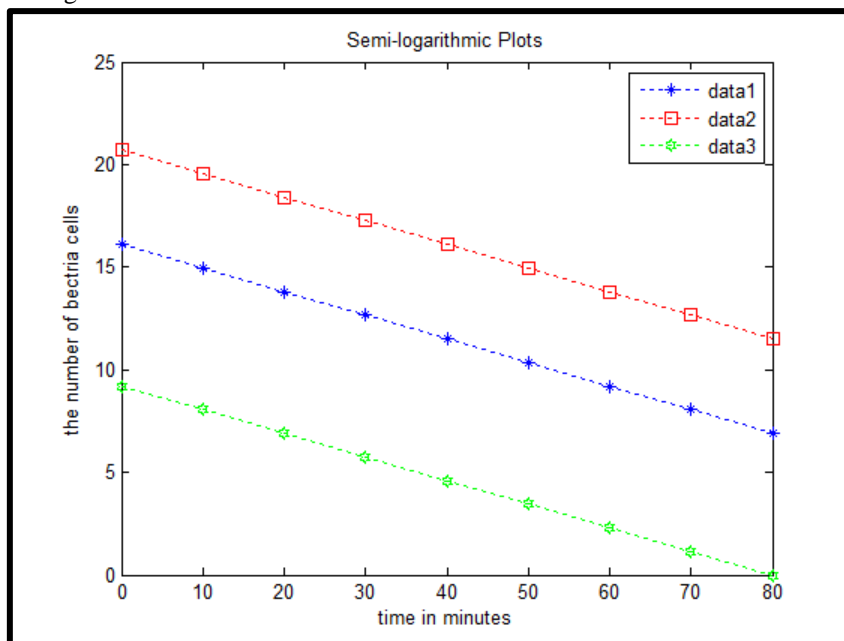


Fig.(6) exponential decay for number of bacteria cells

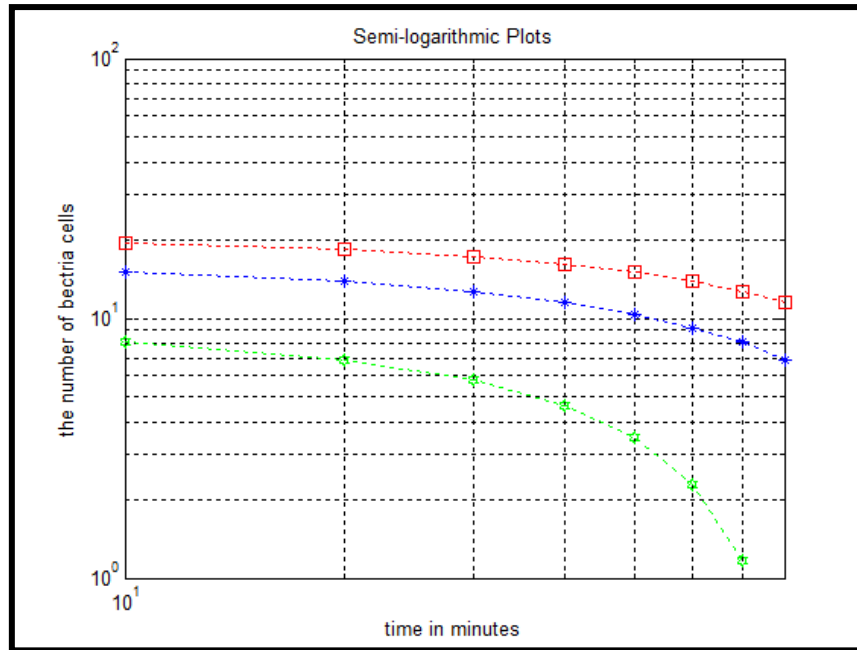


Fig.(7) exponential decay for number of bacteria cells by using paper of graph semi-logarithmic plots

Discussion:

The outcomes effectly affect those studies which depend on the discovery time of bacterial populaces as characterized by the interim slipped by from the vaccination to the time when the populace achieves a specific level. Accepting immaculate exponential ceasing to exist (vanishing) until the season of identification, went before by the slack time frame just, the appropriation of the recognition times mirror the dissemination of the populace slack times, which is increasingly scattered. The recognition time, has higher expected quality and fluctuation, which has evident results, for instance, while evaluating the time until specific indications show up brought about by rot microorganisms in the wake of including anti-infection.

For semi-logarithmic relapse models, we call the "ln tallies v. time" bend the populace bend. At the point when the phones' surroundings all of a sudden gets to be inactivating, (for example, including anti-infection), the "opportunity to death" for a solitary cell will be known as the individual survival time as it show in Fig.(5).

This expression, be that as it may, is joined just for $y=ba^x$; (when $a<1$), and the survival bend portrays an immaculate exponential demise, with $y=9.210e(0.1220)t$ (straight line through the birthplace). On the off chance that $b_2=9.210$ then the expression can't be concurrent, so however the slant of the survival bend is focalized to zero in the wake of including the third anti-microbial as it show in Fig.(6) and Fig. (7).

It ought to be additionally conceivable to estimate the dispersion of the slack times of individual cells from customary reasonable tally development bends, if there is adequate measure of exact estimations before the exponential stage.

In this way, however semi-logarithmic relapse models gives a hypothetical foundation to make identicalness between development bends and slack conveyances, the standard bend fitting techniques, numerically, are not sufficiently strong on normally measured development information. The customary feasible check bends are not reasonable to recognize what sort of conveyance is trailed by the individual cells. New methodologies are fundamental in the estimation systems, as well (picture investigation, stream cytometry, and so forth), keeping in mind the end goal to study singular cell energy.

In any case, three noteworthy purposes of distinction must be considered (Baranyi et al, 1994).

The bacterial fixation locale of concern (for the most part 1 - 109 cells/gram) in ailment microbiology is much more extensive than the focus area that biotechnologists are occupied with. It might be improper subsequently to extrapolate the scientific models connected in biotechnology. Another numerical outcome of the expansive district of interest is that, for measurable/computational cause, it is not the cell fixation but rather its sine-logarithm, which is the favored variable to be displayed.

Mathematical modelling techniques

unavoidably penetrate all sciences, for example, illness microbiology, that was a simply enlightening and experimental science simply a few decades back.

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