HIGH FREQUENCY LIMITS IN PERIODICITY SEARCH FROM IRREGULARLY SPACED DATA

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Abstract. Notions and limits from standard time series analysis must be modified when treating series which are measured irregularly and contain long gaps. Classical Nyquist criterion to estimate frequency range which is potentially recoverable must be modified to handle this more complex situation. When basic exposition of the modified criterion is given in earlier papers, some minor problems and caveats are treated here. Using simple combinatorial arguments we show that for small sample sizes the modified Nyquist limit may overestimate the obtainable frequency range. On the other hand we will demonstrate that very high Nyquist limit values which are typical to irregularly sampled data can often be taken seriously and using proper observational techniques the frequency ranges for “time spectroscopy” can be significantly widened.

Key words: Methods: data analysis – numerical – statistical

1. INTRODUCTION

It is very often the case that observed time series seems to be variable but physical origin of the variability is unknown. The first thing what astronomer does in this case is to compute standard power spectrum or similar statistic for a wide range of trial frequencies and look for possible peaks. But how far along frequency axis one should go with this analysis? Typical answers can be found e.g. in Kurtz (1983), Eyer & Bartholdi (1999) and Koen (2006). Below we look at this question from different angles trying to clarify some theoretical aspects which are untreated till now. Some of our observations show that proposed schemes to estimate upper reasonable frequency limits can be misleadingly high. However, in most cases one tends to underestimate the detection potential of precisely timed and truly randomly spaced measurement series.

2. EQUALLY SPACED DATA

In traditional Fourier analysis the highest frequency that can be extracted from equally spaced and continuously monitored data is \( s_{Nyq} = \frac{1}{2\Delta t} \) (where \( \Delta t \) is sampling step in time) - so called Nyquist frequency (in the context of astronomy, see e.g. Kurtz 1983). This fact can be proved using different analytical tools. For our purposes the shortest route to the result comes from observation that regularly spaced sampling can be looked upon as a multiplication of continuous
input function with appropriately scaled in time III function (see Bracewell 2000):

\[
\text{III}(t) = \sum_{n=-\infty}^{\infty} \delta(t-n).
\]

To the multiplication in time domain corresponds convolution in Fourier domain. The Fourier transform of a III function is also a III function. As a result, we get a sum of periodically (with period \(1/\Delta t\)) repeated original spectra. If the frequency spectrum \(F(s)\) of original data \(f(t)\) is bandlimited by the Nyquist frequency \((F(s) = 0 \text{ when } |s| > 1/(2\Delta t))\) then its full recovery is possible. Otherwise different shifted replicas of the original spectrum overlap and exact recovery is impossible.

\textbf{Fig. 1.} Fourier transforms of \(f(t) = 0.5 \cos(2\pi t \times 0.81) + \cos(2\pi t \times 0.3)\). For lower spectrum \(\Delta t = 1\), for upper spectrum \(\Delta t = 0.25\). Nyquist limits are depicted using thick lines.

This is well illustrated on Figure 1 where two Fourier transforms of the input signal \(f(t) = 0.5 \cos(2\pi t \times 0.81) + \cos(2\pi t \times 0.3)\) are depicted. In upper panel the sampling step \(\Delta t = 0.25\) is short enough to reproduce both frequencies correctly. Spurious peaks start to occur only for frequencies which are higher than Nyquist frequency (in this particular case \(s_{Nyq} = 2\)). In lower panel the transform is computed from data with time step \(\Delta t = 1\). Now the replicas of the original spectra are spaced more densely and inside of the Nyquist limits we can see correct peak at frequency \(s = 0.3\) and spurious peak at frequency \(s = -0.81 + 1 = 0.19\). This kind of overlapping of shifted replicas of the spectrum is generally known as aliasing.

Very often aliasing effect is misinterpreted in a certain way. Normally it is assumed that all non-zero energy of oscillations is situated in the first replica around zero frequency. But this need not be the case. If the actual frequency of the input process is much higher than the Nyquist frequency then its aliases can show up at lower frequencies. From the convolution theorem follows only periodic replication of the spectral fragments, not the actual position of the true
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replica. In some cases this observation allows to observe high frequency phenomena with apparatus which samples input data at significantly lower rate from that required by the Nyquist theorem. The physically relevant frequency from the set of periodically spaced candidates (with step in frequency $\Delta f$) can then be selected using some additional criteria.

3. IRREGULARLY SPACED DATA

In the case of data sets where time moments are measured irregularly the Fourier convolution theorem can be similarly applied (see Deeming 1975). Instead of the III function we can now introduce data window function $w(t)$:

$$w(t) = \sum_{n=1}^{N} \delta(t - t_n),$$

for time point sequence $t_n, n = 1, \ldots, N$. The Fourier image of the sampled data can now be described as a convolution of the real continuous data spectrum $F(s)$ with the Fourier transformed data window function $W(s)$. Typical window function is depicted on Figure 2.

**Fig. 2.** Typical Fourier transformed data window function $W(s)$. Nyquist limits are depicted using thick lines.

The strongest peak is at zero frequency as it should be. But instead of periodically occurring side peaks we can now see only random peaks with significantly lesser amplitude. The important point for a following discussion is that fluctuations inside the Nyquist limits are not very much different if to compare with fluctuations far away.

Now we have a reasonable question - how to put correct limits to frequency search ranges if our input data is randomly sampled?
There are essentially three methods discussed till now, each one based on different heuristic deliberations.

First, it is possible to look at irregularly spaced data set as a disturbed version of the regularly spaced data. Consequently the mean time step is calculated $\Delta t = (t_N - t_1)/(n - 1)$ and corresponding Nyquist limit is calculated using this value. This view is supported also by statistical argument - if we compute spectra from $N$ statistically independent input points then there cannot be more than $N/2$ statistically independent spectrum points. (Another half of the input statistical information is spent on negative frequencies or phases). However, simple trial calculations show that so calculated limits are too conservative and useful information can be lost (see Press et al 2007).

Then it is argued that in the case of irregularly spaced data distances between some consecutive time points are significantly shorter than mean time step and this allows define the Nyquist frequency using minimal time difference between two consecutive observations. Again the numerical experiments (see Eyer & Bartholdi 1999) show that this is not the case and corresponding criterion is too conservative.

The third method (see Eyer & Bartholdi 1999, Koen 2006) takes off from the observation that actual times of observation are always of limited precision and correspondingly their values form a subset of a certain fixed step grid. The ultimate Nyquist limit then can be computed from time step of this grid. The limit can be made more sharp by checking phase distributions for the multiples of the minimal step (check Eq. 7 of the Koen 2006).

4. RETURN ARGUMENT

The third method to estimate the Nyquist frequency is certainly valid, but it contains one important hidden caveat.

Let the Nyquist frequency computed for a particular data set be $s_{Nyq}$. If we want to cover in our analysis frequency range $(0, s_{Nyq})$ with proper frequency resolution we need check at least $K = s_{Nyq}(t_N - t_1)$ trial frequencies (proper step along frequency axis is $\Delta s = (t_N - t_1)/K$). For a extensive set of time point arguments so computed $K$ can be quite large. It is not ruled out that the number of essentially different phase configurations exhausts itself earlier than $K$ trials are computed.

To illustrate this we look first at one old, but still popular method of period seeking, the so called Lafler-Kinman (1965) method. In this method for each trial frequency $s$ phases

$$\phi(t_n, s) = s \times t_n - [s \times t_n], n = 1, \ldots, N$$

are computed (squared brackets denote taking of integer part). Then data points $m(t_n), n = 1, \ldots, N$ are sorted according to the phases. For sorted values (say, magnitudes) the final frequency dependent statistic is computed

$$\theta(s) = \frac{\sum (m_i - m_{i+1})^2}{A}$$

where $m_i$-s are just sorted values for increasing phases, summation takes into account cyclicity ($m_{N+1} = m_1$) and $A$ is certain frequency independent scaling
factor. The minima of the \( \theta(s) \) function indicate then most probable frequencies (it is kind of inverted power spectrum). Very often the methods based on this statistic and similar to that are called as “string length” methods.

For us the important point is that actual frequency dependent phases do not enter into final statistic, their effect is accounted for only through generated sort orders. But there are only \( N! \) different possible orderings (permutations). Consequently, if there is, say only 5 observations in our data set, then there is just 120 different possible permutations and possible \( \theta(s) \) values. It is guaranteed that \( \theta(s) \) values start to reoccur and there is no way to distinguish between them. However for high precision observations the formal Nyquist range can contain much more trial frequencies! As we see the low number of observations themselves puts a certain limit to the number of frequencies which can be inspected using this kind of statistic. Or to put it in another way around the phase configurations start to return before all \( K \) trial frequencies are checked through.

Of course, it can be said that Lafler-Kinman statistic is of very peculiar kind and we just can avoid it. Unfortunately this is not the case. For other statistics it is quite easy to involve similar arguments.

For instance in methods where statistics are computed using binning of phases (see for instance Jurkevich 1971 or Schwarzenberg-Czerny 1989) there can be only \( M^N \) essentially different phase configurations (for each from \( N \) phases we can assign arbitrary bin from \( M \) bins available). For small values of \( N \) and \( M \) it can then happen that \( K > M^N \) and statistic values computed from different phase configurations start to return.

For statistics which depend continuously from phases (standard Fourier spectrum, Lomb-Scargle method etc) situation is more complicated. In principle for a set of precisely measured incommensurable time points the phase configurations and computed from them statistic values never return exactly. But if we fix a certain level of precision it is always possible to estimate approximate return times.

If we imagine frequency dependent phases as points on the circle which rotate with incommensurable speeds then this system will return arbitrary close to whatever state after certain time. This so called Poincaré cycle return time can be approximately estimated (see e.g. Kac 1947) and in our context it scales as \( N^N \).

The heuristic argument behind this value is simple. We can divide the full range of phases \([0, 1)\) into \( M \) bins and approximate continuous base functions (say cos and sin) in piecewise constant manner. This allows us to involve return argument for binned case described above. To use all relevant phase information we can select binning scheme with \( N \) bins and here we are - approximately equal statistic values start to return not later than after \( N^N \) trials.

From the first sight restriction imposed by phase configuration return argument is relevant only for data sets with very low number of observations and need not be taken seriously. However, very often we have data sets which contain considerable number of observations but with peculiar distribution where the observing moments tend to be concentrated in small number of densely populated groups. Now the return argument starts to work for a low frequency part of the spectrum. For a periods which are significantly longer than group lengths the groups behave as singular points.
5. INTEGRATION TIME EFFECTS

Real observations of the variable objects are always obtained using certain integration (or exposure) times to obtain reasonable photon counts and from it a measurable output signal. Let us assume that integration time for every single observation is $\delta t$. Then the measurement procedure can be formally described as a sequence of two operations: convolution of the input continuous waveform by rectangle function $\frac{1}{\delta t} \Pi \left( \frac{t}{\delta t} \right)$ followed by the sampling proper. In the time domain to the convolution with a rectangle function corresponds multiplication with sinc function:

$$\text{sinc}(s\delta t) = \frac{\sin(\pi s\delta t)}{\pi s\delta t}$$

in the Fourier domain. This fact introduces additional restriction to the reasonable frequency range in periodicity search. The convolution acts as a low pass filter and higher frequencies are strongly attenuated. Typical frequency limits obtained from this analysis are in the range from $s_{\max} = \frac{1}{2\delta t}$ to $s_{\max} = \frac{1}{3\delta t}$, (see for instance Eyer & Bartholdi 1999).

Normally the side lobes of the sinc functions are ignored and corresponding higher frequencies are left out from analysis. However, in the case when exposure times are sufficiently exactly measured and exposures are spaced in time sufficiently randomly it is possible to find a proper number of observations $N$ for which peaks in frequency space side lobes can be correctly detected. To cover regions where attenuation is very strong (say around zero crossings) we can use different exposition times. This methodology needs further elaboration and will be treated elsewhere.

6. MODIFIED NYQUIST LIMITS CAN BE TAKEN SERIOUSLY

In previous work (e.g. Press et al 2007, Eyer & Bartholdi 1999 or Koen 2006) the peculiar alias free nature of the irregularly measured samples is demonstrated by computing sample spectra for $2 - 5$ times wider ranges than the “normal” Nyquist ranges (computed from average time step length). From phase configuration return argument above we can conclude that for moderately large values of $N$ phase configurations do not return. Consequently we can safely apply formal Nyquist limits. For precisely measured time point series these limits can be quite high.

To check this in practice we performed some very simple and therefore easily repeatable calculations. First we postulated a standard harmonic model for our data sets:

$$f(t_n) = \cos(2\pi st_n) + E\varepsilon_n, \quad n = 1, \ldots, N$$

where $\varepsilon_n$'s are normally distributed random variables with unit dispersion and $E$ is parameter which controls noise level. The time point sequence $t_n, n = 1, \ldots, N$ was randomly generated with high precision (with numerical grid step $\Delta t \leq 2^{-31}$), using recursive scheme

$$t_n = t_{n-1} + R + 0.25,$$

where $R$ is random computer generated part of the full time step ($0 < R < 1$). The obtained time points were then scaled so that $t_1 = 0.0$ and $t_N = N - 1$ fixing
mean step length at \( \tilde{\Delta}t = 1.0 \) and consequently the standard Nyquist frequency at 0.5. In recursive scheme the fixed part (0.25) of the time step guarantees that in final sequence no step is shorter than 0.2 (the corresponding Nyquist frequency is then 2.5). Formal Nyquist limit which can be computed from grid frequency was approximately \( 2^{10} \) - or in practical terms - infinity. The frequency search range for all runs was fixed at 0.1–500.0 (thousand times the normal range and two hundred times of the range which comes from shortest time distance argument). Step size in frequency was taken as it is normally done \( \Delta s = 1/N \). Corresponding spectra contained then 2000-98980 points (in the range of data point numbers from \( N = 3 \) to \( N = 100 \)). For each trial frequency the standard Lomb-Scargle (Lomb 1976, Scargle 1982) statistic was computed.

To evaluate and compare different spectra we used contrast ratio of the amplitude of the strongest peak in the spectrum to the second strongest. Very often the strongest peaks are evaluated against the mean spectrum level but here we are much more stringent and use highest accidental value from the full spectrum. If the strongest peak in the spectrum was not at correct position the contrast ratio was set to impossible value 0.5. The model oscillation frequency was taken quite high - \( s = 444.444 \).[1]

\[ \text{Fig. 3.} \] Contrast ratios for data sets with different lengths, noiseless case.

As seen from the Figure 3 where one concrete run for a noiseless case \( E = 0 \) is depicted the contrast ratio shows systematic rising trend. The inherent scatter in the contrast ratio values is a result of random distribution of the time points. Because theoretically Lomb-Scargle spectrum has exponential statistical distribution (see Scargle 1982) the occurrence of random strong peaks is quite probable. Nevertheless, it can be seen that already starting from \( N = 17 \) the correct frequency is stably recovered and starting from \( N = 34 \) ratio is permanently higher than 1.5.

[1]The full set of tables from which figures are compiled can be found at http://www.aai.ee/~pelt/soft.htm
The picture of the noisy case with $E = 0.2$ (10% noise level) is quite similar (see Figure 4). The row of the correct detections starts at $N = 26$ and the 1.5 level is achieved at nearly the same level as the noiseless case $N = 43$.

**Fig. 4.** Contrast ratios for data sets with different lengths, 10% noise.

For higher noise levels the picture starts to change. For instance if $E = 0.5$ (25% noise, Figure 5) the ambiguity for short sequences last longer and overall scatter of ratio values is higher. The monotonous exact recovery starts at only
$N = 28$. The 1.5 level stability is achieved at $N = 69$. There is no need to say that these are just particular runs and the results depend on actual random time sequences. However, the general picture is always quite similar.

At the first glance these results seem to be extremely counter intuitive. We compute from, say 50, observations spectra which contain $\approx 50000$ points. In fixed time step situation we can have only 25 statistically independent frequencies below the Nyquist frequency. Now we claim that irregularly sampled scheme can detect true spectrum lines among 50000 different possible positions and quite predictably so. How this can be?

Computed values in the long spectrum for irregularly sampled data are statistically strongly correlated. If we return to our points on circle analogue, then this kind of deterministic systems show also very long (essentially infinite) correlations. Nevertheless the states where all points occur in a certain restricted position are extremely rare (you can compare the situation with molecules in the air). This shows that estimation of the power spectrum (statistical analysis) significantly differs from periodicity seeking (kind of harmonic analysis). In the first case we do not assume that computed spectrum has any prescribed form. Then it is quite clear that we cannot properly estimate more than $N$ (or $N/2$ in power terms) spectrum values from $N$ data points. In the second case we assume that the spectrum contains only one or small number of peaks and our task is to find the exact positions of these peaks. As we demonstrated earlier the frequency range for such analysis is limited by possible return of phase configurations or by the modified Nyquist limit.

7. CONCLUSIONS

From analysis above we can make following conclusions:

- When Galileo used pulse counts and pendula for timing in his experiments and the obtained precision was certainly not better than around one tenth of the second, then modern clocking allows time measurements whose precision can exceed $10^{-16}$ seconds and in the near future even more. This allows to widen frequency range for “time spectrometers” significantly. Even when concrete experiment is seeking for phenomena which show up at lower frequencies, it is always reasonable to record timing data with highest precision available. It is never ruled out that certain follow-up analysis of archival data can make this information usable.

- It is always reasonable to keep integration times with fixed and precisely measured length. For future archival work it is also important to give exact positions of published time points against integration intervals (start, exact middle point or end). This allows properly to combine the data with future observations.

- When analyzing already measured data it is reasonable to compute observing windows for a wide range frequencies to check for the possible peaks which occur not only because of some inherent periodic gaps in timing points but also because of possibility of returning phase configurations. This is especially true for data which consist of small number densely populated short fragments.
In some experiments the maximum time sampling frequency is limited by technical set up (dead times, recording lags etc). It is then always possible to widen frequency range for periodicity search by randomizing measurement moments. If the fixed step scheme is unavoidable, even then we can search for higher frequency aliases at lower parts of spectra. For proper final frequency localization some physical insight can be applied or if possible - prefiltering of the data.

We hope that our considerations help to dispel some common misunderstandings and point to new perspectives of high frequency range experiments which can be of great astrophysical interest.

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