



Arima Model to Predict the Prevalence of Diabetes Type 1 and Type 2 Patients: A Case Study of Jos University Teaching Hospital

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Authors' contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

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Abstract

Diabetes Mellitus is a huge burden for human health, increasing number of patient is likely to result in rising demand for the medical emergencies. Due to limited number of hospitals with standard laboratory test kits to differentiate between type 1 and type 2 diabetes it is important to forecast the future incidences and prepare with proper resource planning. The monthly number of Diabetes patients obtained from Jos University Teaching Hospital is fitted by autoregressive integrated moving average (ARIMA) model. Dataset starting

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from January, 2010 to December,2020. Using ARIMA, several models were evaluated based on the Bayesian Information Criterion (BIC) and Ljung-Box Q statistics. ARIMA(3, 1, 1) is found to be better and used to describe and predict the future trends of Diabetes type 1 and ARIMA(1,1,1) is a better model to predict the future prevalence of diabetes type 2. Therefore, the proposed model will help in the appropriate planning and allocation of resources for emergencies.

Keywords: Diabetes; mellitus; hospital; ARIMA; statistics; forecasting; BIC.

1 Introduction

“Diabetes mellitus is chronic syndrome associated with long term complications to the brain, kidney, and the heart. There is destruction and loss of the β -cells of the pancreas causing insulin deficiency; it may also result from abnormalities arising from resistance to insulin. Symptoms of hyperglycemia include polydipsia, polyphagia polyuria, blurred vision, weight loss, generalized pruritus, neuropathy, retinopathy, etc. Life threatening consequences of uncontrolled diabetes include diabetes-ketoacidosis, lactic acidosis and hyperosmolar non-ketotic state” [1]. Diabetes is preceded by impaired fasting glucose (IFG) resulting in a pre-diabetic state which can exist undetected for many years, [2], causing irreversible damage to vital organs. Pre-diabetes is a practical term referring to Impaired Fasting Glucose (IFG), impaired glucose tolerance [3] or a glycosylated hemoglobin (A1c) of 6.0 to 6.4%, each of which places individuals at high risk of developing diabetes and its complications. “The World Health Organization criteria for diagnosing pre-diabetes are fasting plasma glucose level of between 6.1 and 6.9 mmol/l. A fasting plasma glucose level 7.0 mmol/l or more meets the criteria for the diagnosis of diabetes. Fasting value for venous and capillary plasma glucose are identical” [4].

“*Type1 diabetes* occur as a results of auto-immune β -cell destruction in thePancreas characterized by a total absence of insulin production. Type 1 diabetes is responsible for 5% to 10% of all cases of diabetes. Associated risk factors include autoimmune, genetic, and environmental factors”, [5].

“*Type2 diabetes* can be linked to be accounting for around 90 per cent of all cases, it is a chronic metabolic disorder, in which the body is unable utilize glucose from food because of the inability of the pancreas to produce insulin or produces insufficient insulin, or the insulin itself is inactive” [6].

“*Gestational diabetes* could be described as a form of glucose intolerance that affects some women during pregnancy. This kind of diabetes is triggered during pregnancy. Most gestational diabetes is resolved naturally after delivery, but 5-10 percent of women affected during pregnancy are later found to have diabetes, especially Type 2, after pregnancy”.

“*Pre-diabetes* is described as a precursor condition to diabetes in which a person is experiencing elevated blood glucose levels but not yet up to diagnostic criteria for diabetes. People with pre-diabetes normally suffer from impaired fasting glucose or impaired glucose tolerance, or both. Prevention of diabetes mellitus disease requires” [6].

“There is an increasing prevalence of diabetes and pre-diabetes worldwide” [7]. “Over 5 million people suffer from the disease in Africa and the number is expected to skyrocket to 15 million by 2025”, [7]. “In Nigeria the prevalence varies from 0.65% in rural Mangu village to 11.0% in urban Lagos” [8]. “With the incidence of diabetes in Africa, diabetic complications are also expected to rise proportionately” [9,10]. “This will undoubtedly pose serious health and economic problems. The disease affects many people under the age of 64 years in Africa as compared to the developed world where it affects many people over the age of 64 years” [9]. “In Nigeria the National prevalence of diabetes was 2.2%” [8]. In South Eastern Nigeria the overall prevalence of diabetes was 10.51% [11], whereas in South Western Nigeria the prevalence of diabetes ranges from 4.76% in Ile-Ife, Osun State to 11.0% in Lagos [8,7]. “Also 0.8% of diabetes mellitus, and 2.2% of Impaired Glucose Intolerance in Ibadan” [12]. This was comparable to WHO reported a prevalence of 2.8% in Ibadan [13], and 6.8% in Port Harcourt, Nigeria [14]. In 2004, a survey in Jos [14] reported a prevalence of 10.3%. [14] reported “a prevalence of 2.2% in Port Harcourt in 2003”. A prevalence of 4.7% was reported by [15] which was higher than the national prevalence of 2.2% reported by [7]. A review of studies on the prevalence of diabetes in adults in Africa [16] demonstrated a rising prevalence across the continent.

“Time series analysis is one of the quantitative methods which can effectively predict the future incidence of communicable diseases and epidemiological trends using previously observed data and time variables”, [17]. “This analysis deals with time dependent variables with an advantage of being not necessary to consider the influence of intricate factors” [18,19].

“Time series methods have been widely used to analyzed infectious diseases’ surveillance data in recent decades, including data for sexually transmitted diseases. Different time series models were used to forecast the epidemic behaviour in previous study” [20]. “For example, decomposition methods were used to forecast nine notifiable infectious diseases in China” [21]. Autoregressive Integrated Moving Average Models (ARIMA) are widely applied in infection time series modelling including tuberculosis [22], typhoid fever [20], gonorrhoea [23] and hepatitis [24]. “The Autoregressive conditional heteroscedasticity and generalized autoregressive conditional heteroscedasticity models have been used to investigate the risk factors associated with syphilis”, [25]. Seasonal Autoregressive Integrated Moving Average (SARIMA) model has been increasingly favoured and successfully used in the prediction of communicable diseases, such as dengue [26], tuberculosis [27], mumps [19] and others [27-29]. “According to the International Diabetes Federation (IDF), there were 463 million adults with diabetes worldwide in 2019, an average of 1 in 11 adults. Furthermore, there were 4.2 million individuals who died from diabetes and its complications, accounting for about 11.3% of all global deaths” [30]. “Another study reported that there were 67.9 million disability-adjusted life years (DALYs) attributable to diabetes globally in 2017. In the global ranking of DALYs caused by disease, DALYs caused by diabetes ranks ninth for females and eighth for males” [31–33]. “In addition, alongside changes in dietary structure and irregular living habits in modern society over recent years, the incidence, morbidity, and mortality of incidence has shown an increasing trend year by year” [34]. “According to the tenth edition of the Diabetes Map released by the IDF, there were 537 million patients in diabetes globally in 2021, accounting for 10.5% of the global population. Compared with data reported in the ninth edition of the Diabetes Map, the prevalence rate of diabetes increased by 12.9% overall. The number of diabetic patients is expected to reach 578 million by 2030, and estimated to reach 783 million worldwide by 2045” [35]. “The control and treatment of this disease requires substantial medical resources” [36]. “Other studies have shown that DALYs increased by 30.0% from 2007 to 2017, years of life lost (YLL) increased by 29.9%, and years lived with disability (YLD) increased by 30.1%” [31–33]. “Moreover, the economic expenditure related to diabetes globally was substantial, reaching 1054 billion dollars by 2045” [37]. “The widespread prevalence of diabetes has brought huge burden to the social and economic development of the world, and the situation is not optimistic. Diabetes has become a global public health problem. How we can effectively treat diabetes, slow down the occurrence of complications, and greatly improve the life quality of patients, have become critical medical problems that needed to be solved urgently” [37].

“Over recent years, numerous models have been used to assess and predict the disease burden associated with diabetes. The main model used to assess disease burden is the linear regression model” [38]. For example, the Global Burden of Disease (GBD) 2019 Dementia Prediction Collaborative Group [38] used a linear regression model to investigate changes in the global prevalence of dementia. Li [39] used “a linear regression model to evaluate the disease burden associated with esophageal cancer. Many models have been used for disease burden; the most common models are the age-period-cohort (APC) model and the time series model”. Ji [40] used “the APC model to predict the incidence of hepatitis B in males and females of specific age groups”. In another study, Akita [41] used “the APC model to predict the mortality associated with hepatocellular carcinoma and recapitulated the observed mortality”. Zheng [42] explored “the feasibility of using the auto-regressive integrated moving average model (ARIMA) and the Elman neural network model to predict the incidence of hepatitis B”. Ceylan [43] used “the ARIMA model to predict the prevalence trend of COVID-19 in the three countries most affected by COVID-19 in Europe, including Spain, Italy, and France”. In another study, Fu [44] used “the exponential smoothing model (ES) model to predict the incidence of acute upper gastrointestinal bleeding. However, no previous study has applied these models to assess and predict the disease burden related to diabetes. This study aimed to complement and refine this aspect of analysis by applying these common and reliable models”.

“Accurate disease burden data is an important basis for scientific development and the timely adjustment of health policies and strategies. Such data can contribute to the development of clear prevention and control priorities, and help to evaluate the efficacy of measures. Some studies have examined the disease burden associated with diabetes in specific regions. These studies found that although the mortality rate associated with

diabetes was declining, the burden of diabetes was still increasing“ [45]. “Other studies showed that the impact of diabetes on cancer was increasing” [46].

Different models have been used in the analysis of diabetes but there are few of such in the study areas and none has considered the types of diabetes. Hence, this research is aimed at developing a model that will forecast the prevalence of diabetes type 1 and type 2 using secondary data by selecting the best fit models to predict the trends of diabetes type1 and type2 diabetes and forecasting diabetes type 1 and type 2 in the next 5 years using the selected models.

2 Materials and Methods

Autoregressive (AR) Model: An Autoregressive (AR) model of order P satisfied the following

$$\begin{aligned}
 x_t &= \mu + \sum_{k=1}^p \phi_{kk}x_{t-k} + \varepsilon_t \\
 x_t &= \mu + \phi_{11}x_{t-1} + \phi_{22}x_{t-2} + \dots + \phi_{pp}x_{t-p} + \varepsilon_t \quad \dots \dots \dots (3.1)
 \end{aligned}$$

For t=0 where $\varepsilon_t = 0$

For n> 0 where the error term $\varepsilon_t > 0$, is a series of independently, identically distributed (i.i.d) random variables and assumed to be normally distributed and μ is some constant. The p denotes the order of autoregressive model, defining how many previous values the current value is related to. The model is called autoregressive because the series is regressed on to past values of itself. The error term ε_t in equation (3.1) refers to the noise in the time series. Above, the errors were said to be independently identically distributed. Commonly, they are also assumed to have a normal distribution with mean zero and variance σ^2 . For the model in equation (3.1) to be of use in practice, the estimator must be able to estimate the value of ϕ_k

Moving Average (Ma) Model: An Autoregressive moving Average of order q is given as

$$\begin{aligned}
 x_t &= \mu + \sum_{j=1}^q \theta_{jj}x_{t-j} + \varepsilon_t \\
 x_t &= \mu + \theta_{11}x_{t-1} + \theta_{22}x_{t-2} + \dots + \theta_{pp}x_{t-q} + \varepsilon_t \quad \dots \dots \dots 3.2
 \end{aligned}$$

For t=0 where $\varepsilon_t = 0$

The error (or noise) term in this equation is the one step ahead forecasting error which can be expressed as a function of previous forecasting errors. It shows that MA (q) models make forecast based on the error made in the past, and so one can learn from the error made in the past to improve current forecast.

Autoregressive Moving Average (ARMA) Models: This is Autoregressive Component of order p and Moving Average Component of order q. In a situation where AR (q) and MA(p) were not able to solve the case at hand, a combination of the two produces another interesting model known as Autoregressive moving average (ARMA)(p, q) model where p is the number autoregressive component and q is the number of moving average component. This model is as presented as follows

The Form ARMA (p, q) model is given by the equation

$$\begin{aligned}
 x_t - \phi_{11}x_{t-1} - \phi_{22}x_{t-2} - \phi_{33}x_{t-3} - \dots - \phi_{pp}x_{t-p} &= \varepsilon_t - \theta_{11}x_{t-1} - \theta_{22}x_{t-2} - \theta_{33}x_{t-3} - \dots - \\
 \phi_{pp}x_{t-p} &\dots \dots \dots 3.3
 \end{aligned}$$

Where ϕ 's (phis) are the autoregressive parameters to be estimated, the θ 's (thetas) are the moving average parameters to be estimated the x 's are the original time series values and the ε 's a residuals or errors which are assumed to follow the normal probability distribution.

In order to make the writing of model easier, Box-Jenkins use the backshift operator. The backshift operator B can change the time period t to time period $t - 1$. For example, $BY_t = Y_{t-1}$, $B^2Y_t = Y_{t-2}$ and so on.

So the above model can be rewritten as:

$$(1 - \phi_1 B - \dots - \phi_p B^p)X_t = (1 - \theta_1 B - \dots - \theta_q B^q)\varepsilon_t \quad \dots \dots \dots 3.4$$

Further can be abbreviated as:

$$\phi_p(B)X_t = \theta_q(B)\varepsilon_t$$

where $\phi_p(1 - \phi_1 B - \dots - \phi_p B^p)$ and $\theta_q(1 - \theta_1 B - \dots - \theta_q B^q)$

Autoregressive Integrated Moving Average (ARIMA) Model: “In statistics, ARIMA(pdq) models, sometimes called the Box-Jenkins models after the iterative Box- Jenkins methodology usually used to estimate them, are typically applied to time series data for forecasting. Given a time series of data $X_{t-1}, X_{t-2}, X_2, X_1$, the ARIMA model is a tool for understanding and, perhaps, predicting future values in the series. The model consists of three parts: an Autoregressive (AR) part, a moving Average (MA) part and the differencing part. The model is usually then referred to as the ARIMA (p, d, p) model where p is the order of the Autoregressive part, d is the order of differencing and q is the order of the moving Average part”, [47].

“However, it is impossible to have the data always stationary. Therefore, to make the process we have to do differencing in non-stationary series so the differenced series $(1 - B)^d X_t$ must be add in the process ARMA (p, q). Where the a_t is a sequence of identically distributed uncorrelated deviates, referred to as "white noise. In many situations where differencing is employed, a non-zero constant term will not be required” [48]. For brevity, the equation is generally written as:

$$\phi_p(B)(1 - B)^d X_t = \theta_0 + \theta_q(B)\varepsilon_t \quad \dots \dots \dots 3.6$$

Where

$$X_t = a_t + \sum_{k=1}^p \phi_k X_{t-k} + \varepsilon_t + \sum_{j=1}^q \theta_j \varepsilon_{t-j}$$

and $(1 - B)^d X_t$ is a differenced series and ε_t 's is a sequence of identically distributed uncorrelated deviates, referred to as "white noise".

2.1 Forecasting using Arima model

After describing various time series models, the next issue to our concern is how to select an appropriate model that can produce accurate forecast based on a description of historical pattern in the data and how to determine the optimal model orders. Statisticians George Box and Gwilym Jenkins developed a practical approach to build ARIMA model, which best fit to a given time series and also satisfy the parsimony principle. Their concept has fundamental importance on the area of time series analysis and forecasting. In this section we will consider the Box Jenkins model building techniques, these consist of the following four steps:

Preliminary Transformation: If the data shows characteristics violating the stationarity assumption, then it may be necessary to make a transformation so as to produce a series compatible with the assumption of stationarity. After appropriate transformation, if the sample autocorrelation function appears to be non-stationary, differencing may be carried out.

Model Identification: “if $\{yt\}$ is the stationary series obtained in step 1, the problem at the identification stage is to find the most satisfactory ARIMA (p,q) model to represent $\{yt\}$ ” [49]. Determined the integer parameters (p,q) that governs the underlying process $\{yt\}$ by examining the autocorrelations function (ACF) and partial autocorrelations (PACF) of the stationary series.

Estimation of the model: This deal with estimation of the tentative ARIMA model identified in step 2. In this study, we use the Statistical Package Social Sciences (SPSS 25) software to estimate the coefficient.

Diagnostic Checking: Having chosen a particular ARIMA model, and having estimated its parameters, the adequacy of the model is checked by analyzing the residuals. If the residuals are white noise; we accept the model, else we go to preliminary transformation stage again and start over.

3 Results and Discussion

3.1 Model for type 1 diabetes

Model Identification: The monthly Diabetes type 1 recorded in Jos University Teaching Hospital from 2010-2020 is plotted in the Fig. 1. The data display trends, which indicate that the mean is not stationary. However, it is necessary to check variance stationarity first.

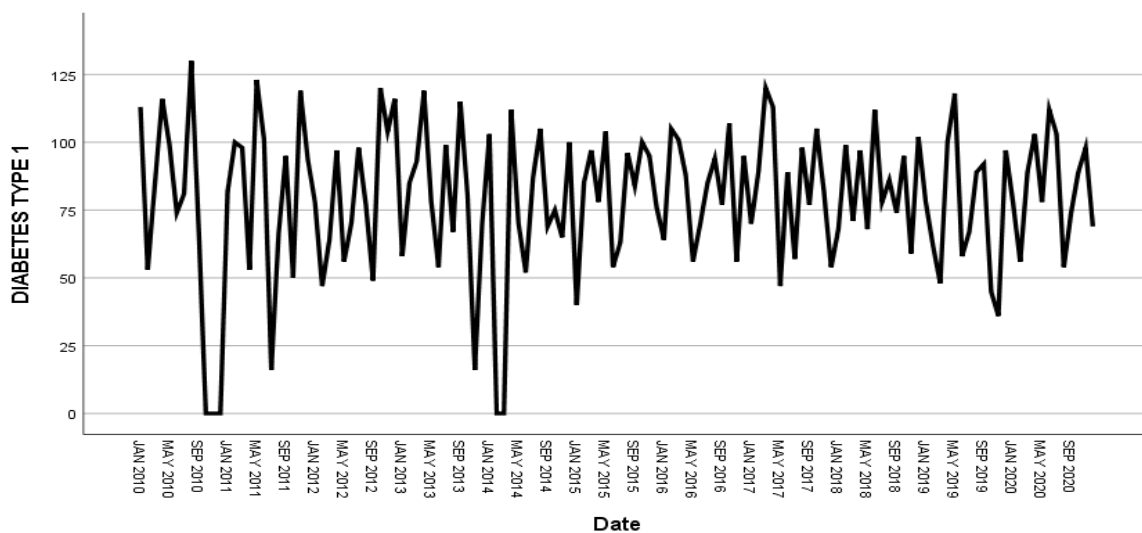


Fig. 1. Monthly number of diabetes type 1 patients from Jos university teaching hospital (January 2010-December 2020)

3.2 Test for stationarity of the variance

It is necessary to check the stationarity of the variance. We can use Shapiro-Wilk and Kolmogorov-Smirnov for the normality test. According to the result providing in Table 1, we can conclude that the variance was not constant (p -value < 0.05). This problem can be corrected by a first-degree order of differentiation.

Further, to make the mean stationary, first order differencing is used. The monthly diabetes type 1 data with first order differencing is plotted in the Fig. 2. Notice that the differenced series is mean stationary and constant variance.

3.3 Graphical representation of autocorrelations

“From the ACF plots below (Fig. 3), the value of its function is less than one. The parameter of any given function is less than one; it means that there is stationarity. Further the plot decays to zero on the both sides of the mean. From the Fig. 3 it can be concluded that an appropriate model is moving average process of order 1” [48].

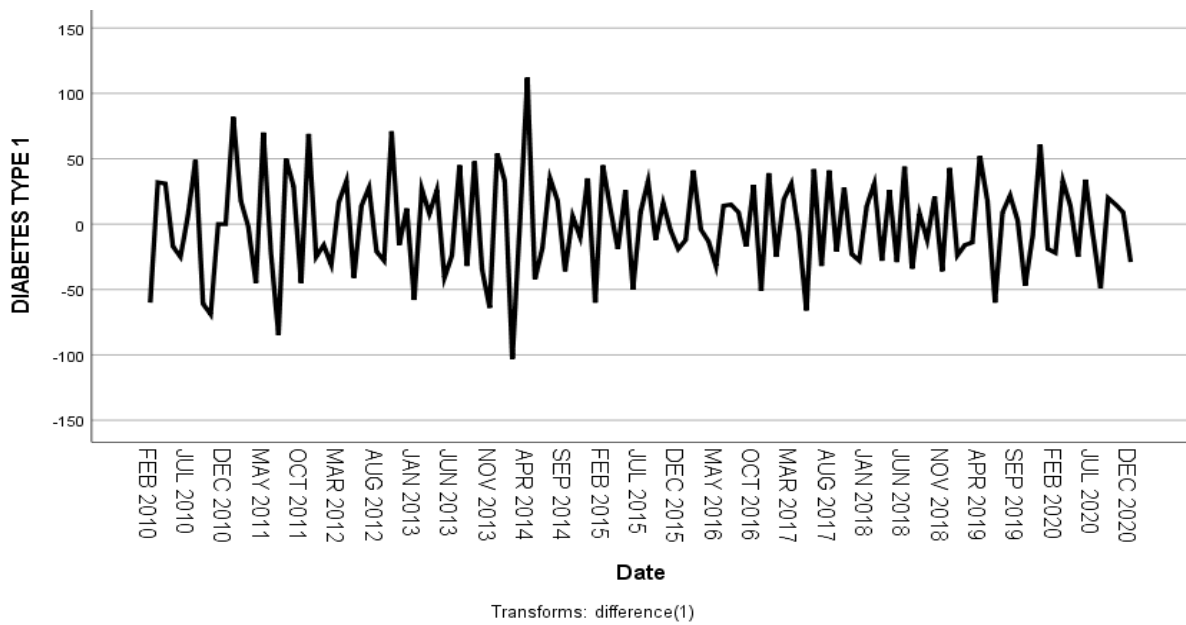


Fig. 2. Time series plot of Diabetes type1patients after the first differencing

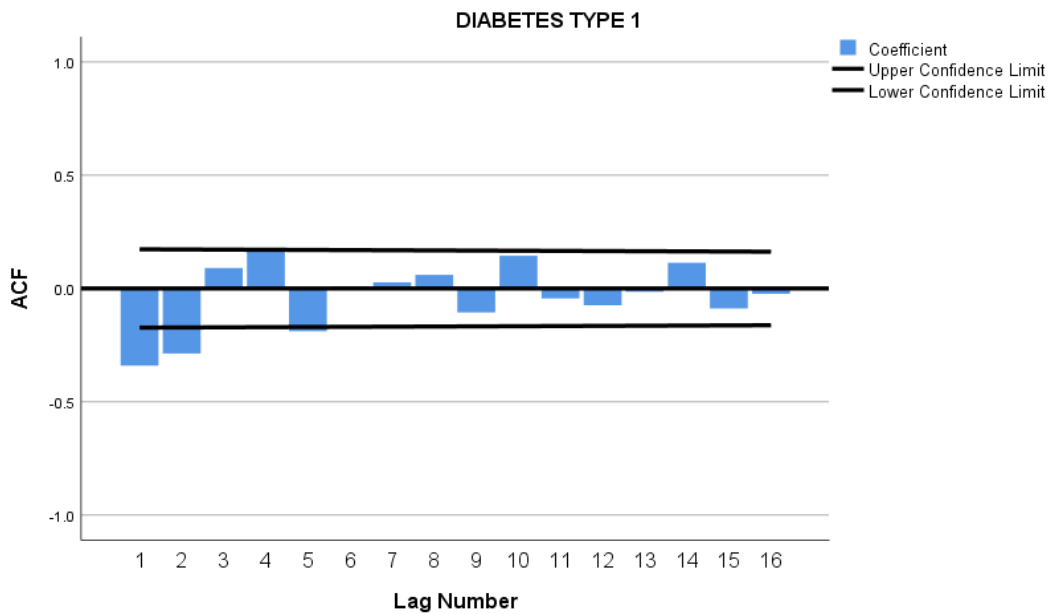


Fig. 3. Sample ACF plots of monthly transformed Diabetes type1 data after first differencing

Observing the plots of ACF and PACF plot above, it could be noticed that the ACF displays the sharper cuts off compared to the PACF. According to the PACF plot in Fig. 4, we use the autoregressive process of order 3, i.e., AR(3).

3.4 Parameter estimation

Four models are chosen and tested to find a model with good fit. These models include ARIMA(1,1,0), ARIMA(2,1,0), ARIMA(3,1,0), and ARIMA(3,1,1)

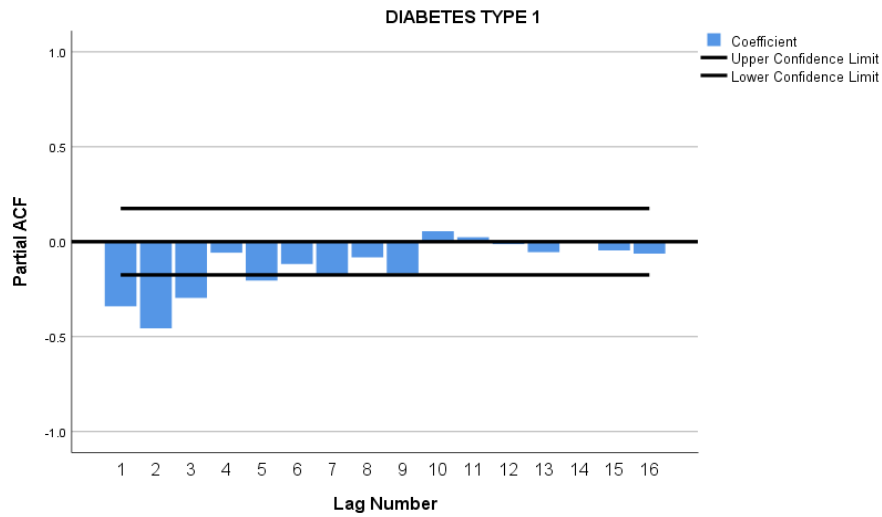


Fig. 4. Sample PACF plot of monthly transformed Diabetes type1 data after first differencing

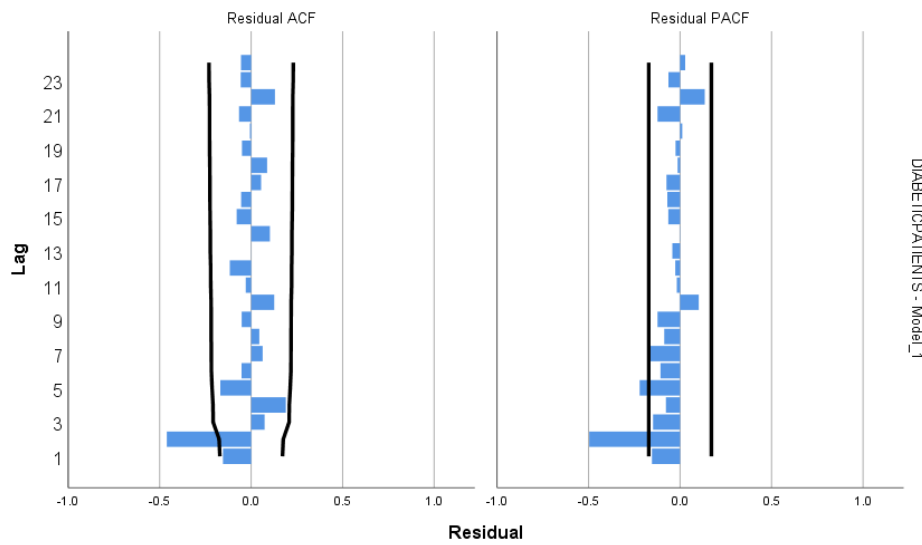


Fig. 5. Residual plots for ACF and PACF after estimating ARIMA (1,1,0)

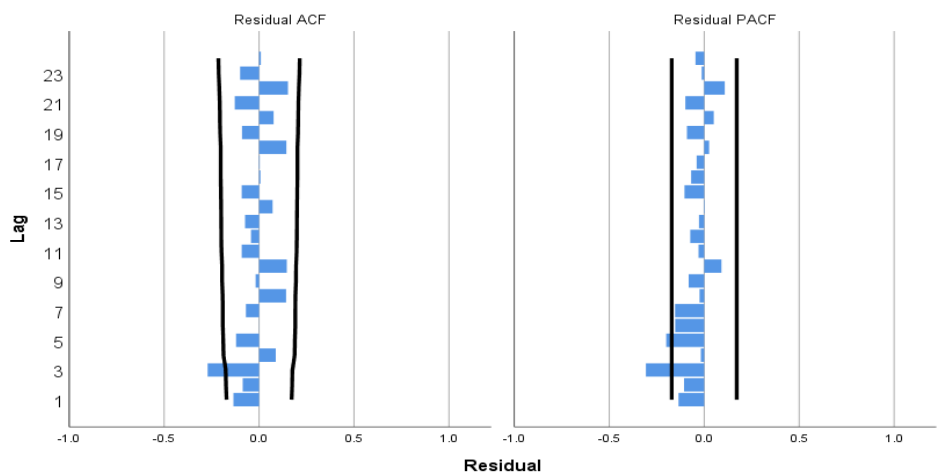


Fig. 6. The residual plots of ACF and PACF after estimating ARIMA (2,1,0)

Table 1. Shows the model summary of ARIMA (1,1,0)

Fit Statistic	Mean	SE	Minimum	Maximum	Model Fit						
					Percentile						
					5	10	25	50	75	90	95
Stationary R-squared	.116	.	.116	.116	.116	.116	.116	.116	.116	.116	.116
R-squared	-.635	.	-.635	-.635	-.635	-.635	-.635	-.635	-.635	-.635	-.635
RMSE	35.012	.	35.012	35.012	35.012	35.012	35.012	35.012	35.012	35.012	35.012
MAPE	42.199	.	42.199	42.199	42.199	42.199	42.199	42.199	42.199	42.199	42.199
MaxAPE	577.529	.	577.529	577.529	577.529	577.529	577.529	577.529	577.529	577.529	577.529
MAE	27.658	.	27.658	27.658	27.658	27.658	27.658	27.658	27.658	27.658	27.658
MaxAE	112.218	.	112.218	112.218	112.218	112.218	112.218	112.218	112.218	112.218	112.218
Normalized BIC	7.186	.	7.186	7.186	7.186	7.186	7.186	7.186	7.186	7.186	7.186

Table 2. ARIMA model parameters of ARIMA (1,1,0)

			Estimate	SE	t	Sig.	
DIABETIC PATIENTS-Model_1	DIABETIC PATIENTS	No Transformation	<u>Constant</u>	-1.162	2.273	-.071	.943
			<u>AR Lag 1</u>	-.346	.083	-4.188	.000
			Difference	1			

Table 3. Shows the model summary of ARIMA(2,1,0)

Fit Statistic	Mean	SE	Minimum	Maximum	Model Fit						
					Percentile						
					5	10	25	50	75	90	95
Stationary R-squared	.301	.	.301	.301	.301	.301	.301	.301	.301	.301	.301
R-squared	-.294	.	-.294	-.294	-.294	-.294	-.294	-.294	-.294	-.294	-.294
RMSE	31.271	.	31.271	31.271	31.271	31.271	31.271	31.271	31.271	31.271	31.271
MAPE	36.591	.	36.591	36.591	36.591	36.591	36.591	36.591	36.591	36.591	36.591
MaxAPE	397.829	.	397.829	397.829	397.829	397.829	397.829	397.829	397.829	397.829	397.829
MAE	25.216	.	25.216	25.216	25.216	25.216	25.216	25.216	25.216	25.216	25.216
MaxAE	82.151	.	82.151	82.151	82.151	82.151	82.151	82.151	82.151	82.151	82.151
Normalized BIC	6.997	.	6.997	6.997	6.997	6.997	6.997	6.997	6.997	6.997	6.997

Table 4. ARIMA Model Parameters ARIMA(2,1,0)

					Estimate	SE	t	Sig.
DIABETIC PATIENTS- Model_1	DIABETIC PATIENTS	No Transformation	Constant		-.077	1.393	-.055	.956
			AR	Lag 1	-.503	.079	-6.371	.000
				Lag 2	-.461	.078	-5.882	.000
			Difference	1				

Table 5. Shows the model summary of ARIMA (3,1,0)

Fit Statistic	Mean	SE	Minimum	Maximum	Model Fit						
					Percentile						
					5	10	25	50	75	90	95
Stationary R-squared	.362	.	.362	.362	.362	.362	.362	.362	.362	.362	.362
R-squared	-.181	.	-.181	-.181	-.181	-.181	-.181	-.181	-.181	-.181	-.181
RMSE	29.991	.	29.991	29.991	29.991	29.991	29.991	29.991	29.991	29.991	29.991
MAPE	33.699	.	33.699	33.699	33.699	33.699	33.699	33.699	33.699	33.699	33.699
MaxAPE	434.673	.	434.673	434.673	434.673	434.673	434.673	434.673	434.673	434.673	434.673
MAE	23.023	.	23.023	23.023	23.023	23.023	23.023	23.023	23.023	23.023	23.023
MaxAE	75.782	.	75.782	75.782	75.782	75.782	75.782	75.782	75.782	75.782	75.782
Normalized BIC	6.951	.	6.951	6.951	6.951	6.951	6.951	6.951	6.951	6.951	6.951

Table 6. ARIMA Model Parameters of ARIMA (3,1,0)

				Estimate	SE	t	Sig.
DIABETIC PATIENTS-Model_1	DIABETIC PATIENTS	No Transformation	Constant	-.077	1.031	-.075	.940
			AR Lag 1	-.639	.085	-7.504	.000
			AR Lag 2	-.610	.086	-7.085	.000
			AR Lag 3	-.298	.085	-3.517	.001
			Difference	1			

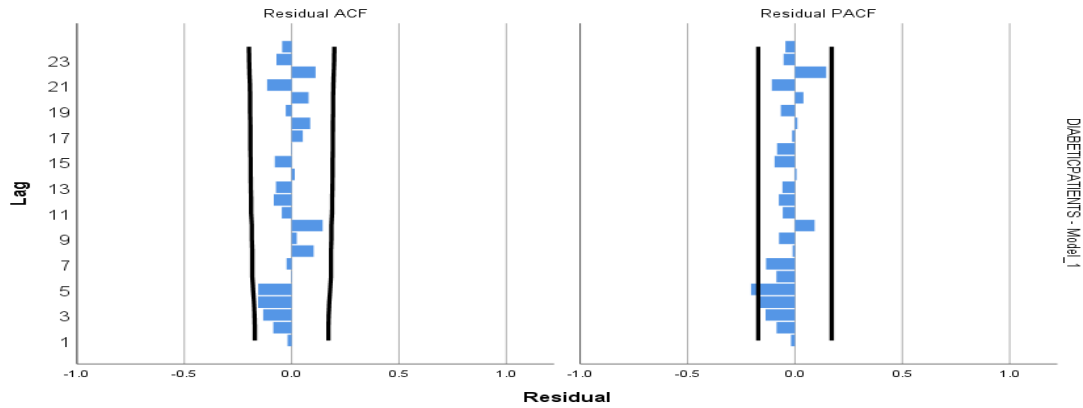


Fig. 7. Shows the residual plots of ACF and PACF after estimating ARIMA(3,1,0)

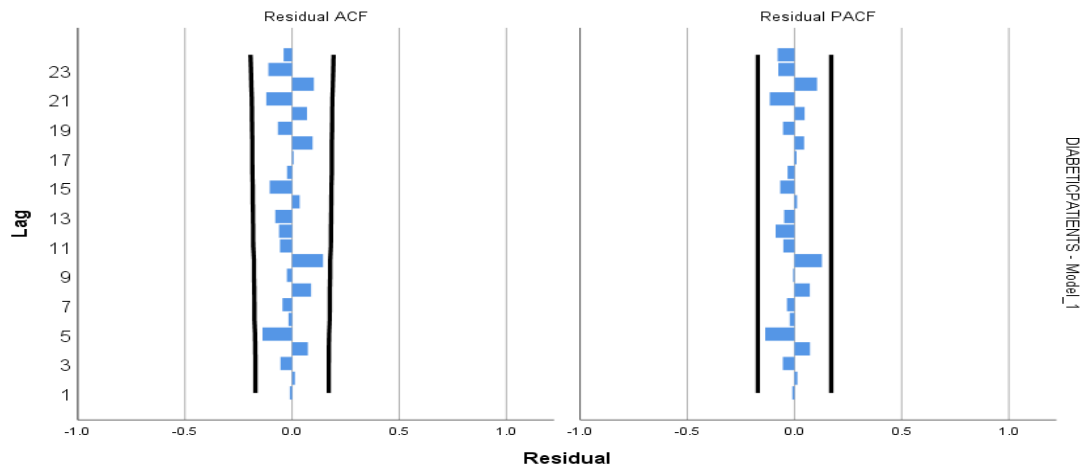


Fig. 8. Residual plots of ACF and PACF after estimating ARIMA (3,1,1)

Based on the Ljung-Box Q statistics, the models with significant p-values are selected and shown in Table 9 below. The resultant BIC providing in Table 10 shows that the most appropriate model to fit diabetes type 1 patients’ data is ARIMA (3, 1, 1).

3.5 Diagnostic checking and parameter estimation

“Diagnostic checking is also known as the verification. It is to do with the testing of the goodness of fit test statistics of a model. We study the ACF and PACF of the residual plot to see it is white noise. If all the autocorrelation and partial correlation are small then the model is working fine but if some of the autocorrelations are large, the values of p or q are adjusted and the model is re-estimated. The checking of residuals and adjusting the values of p and q continues until the resulting residuals contain no additional Structure” [48].

Table 7. Model Summary of ARIMA (3,1,1)

Fit Statistic	Mean	SE	Minimum	Maximum	Model Fit						
					Percentile						
					5	10	25	50	75	90	95
Stationary R-squared	.455	.	.455	.455	.455	.455	.455	.455	.455	.455	.455
R-squared	-.009	.	-.009	-.009	-.009	-.009	-.009	-.009	-.009	-.009	-.009
RMSE	27.832	.	27.832	27.832	27.832	27.832	27.832	27.832	27.832	27.832	27.832
MAPE	29.653	.	29.653	29.653	29.653	29.653	29.653	29.653	29.653	29.653	29.653
MaxAPE	349.071	.	349.071	349.071	349.071	349.071	349.071	349.071	349.071	349.071	349.071
MAE	21.432	.	21.432	21.432	21.432	21.432	21.432	21.432	21.432	21.432	21.432
MaxAE	80.625	.	80.625	80.625	80.625	80.625	80.625	80.625	80.625	80.625	80.625
Normalized BIC	6.838	.	6.838	6.838	6.838	6.838	6.838	6.838	6.838	6.838	6.838

Table 8. ARIMA model parameters of ARIMA(3,1,1)

				Estimate	SE	t	Sig.	
DIABETIC PATIENTS-Model_1	DIABETIC PATIENTS	No Transformation	Constant	.056	.067	.835	.405	
			AR					
				Lag 1	.134	.093	1.443	.152
				Lag 2	-.207	.089	-2.335	.021
				Lag 3	.102	.092	1.109	.269
			Difference	1				
			MA	Lag 1	.998	.570	1.751	.082

Table 9. Competitive models for the monthly diabetes patients based on the Ljung-Box Q statistics test

Models	Q-statistics	P-value
ARIMA(1,1,0)	52.091	.000
ARIMA(2,1,0)	30.689	.015
ARIMA(3,1,0)	19.379	.197
ARIMA(3,1,1)	13.597	.480

Table 10. Competitive models for the monthly diabetes patients

Fit statistic	ARIMA(1,1,0)	ARIMA(2,1,0)	ARIMA(3,1,0)	ARIMA(3,1,1)
Stationary R-square	.116	.301	.362	.455
R-square	-.635	-.294	-.181	-.009
RMSE	35.012	31.271	29.991	27.832
MAPE	42.199	36.591	33.699	29.653
MaxAPE	577.529	397.829	434.673	349.071
MAE	27.658	25.216	23.023	21.432
Normalized BIC	7.186	6.997	6.951	6.838

Form Fig. 8 the sample ACF and PACF of the model shows that the autocorrelations of the residual are all close to zero which mean they are uncorrelated, hence the residual assume mean of zero and constant variance. Finally, the p-value (0.480) for the Ljung-Box statistic clearly exceeds 5% for all lag orders. Thus, the selected model ARIMA (3, 1, 1) satisfies all the model assumptions.

3.6 Forecasting using ARIMA (3, 1, 1)

The forecast values with 95 percent forecast limit of the ARIMA (3, 1, 1) of model for monthly diabetesPatients are shown in Table 16 with standard error, lower and upper limit and its actual forecasted values.

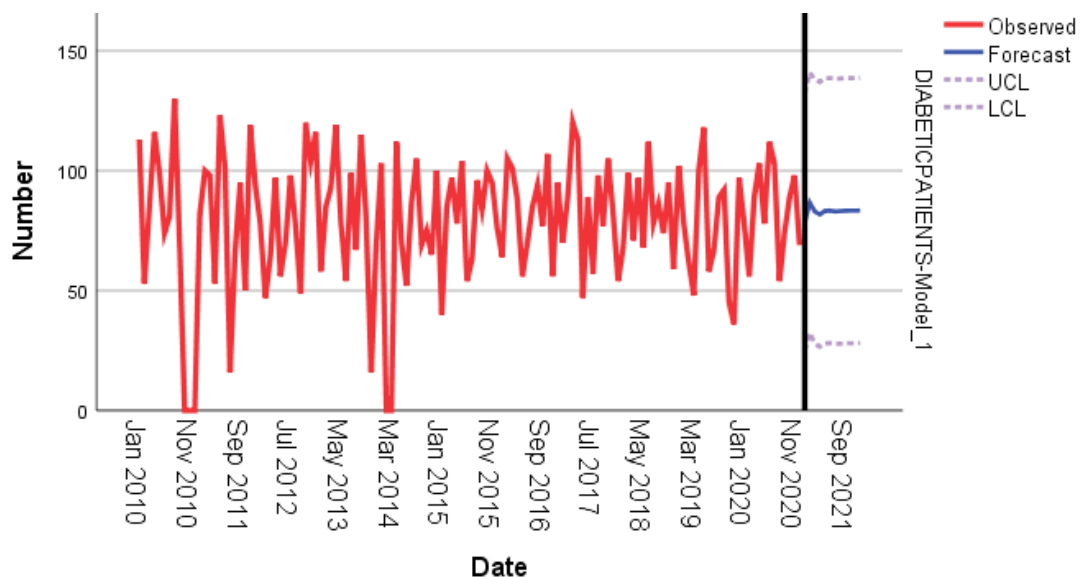


Fig. 9. Forecast chat of ARIMA (3,1,1)

3.7 Model for type 2 diabetes

Model Identification: The monthly Diabetes type 2 recorded in Jos University Teaching Hospital from 2010-2020 is plotted in the Fig. 13. The data plot display trends, which indicates that the mean is not stationary. This can be corrected by first order differentiation.

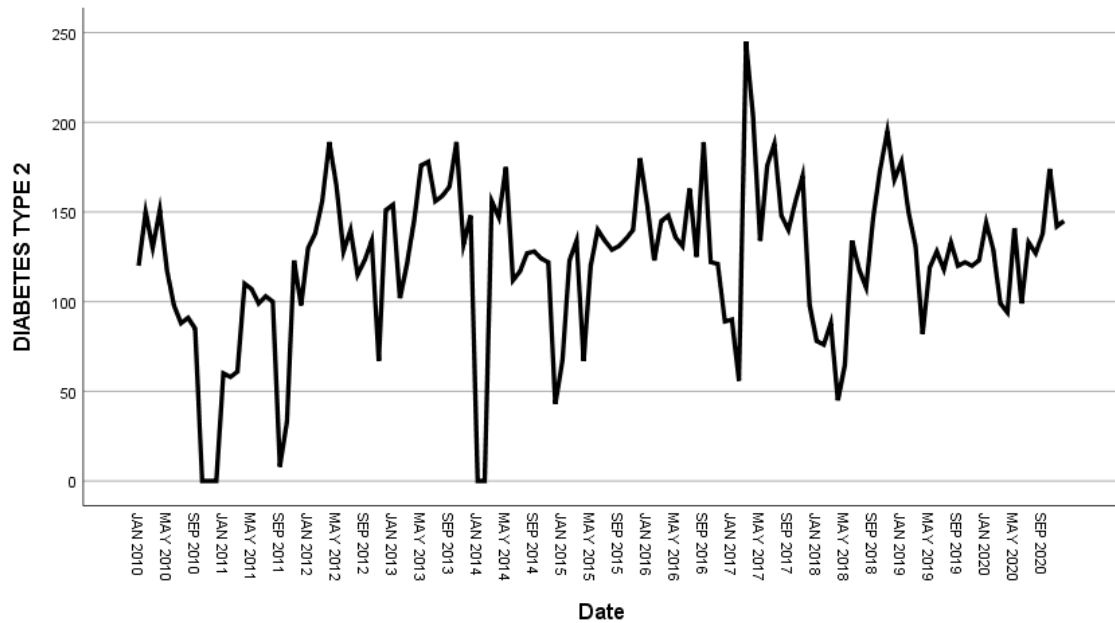


Fig. 10. Monthly number of diabetes type 2 patients from Jos University Teaching Hospital (January 2010- December 2020)

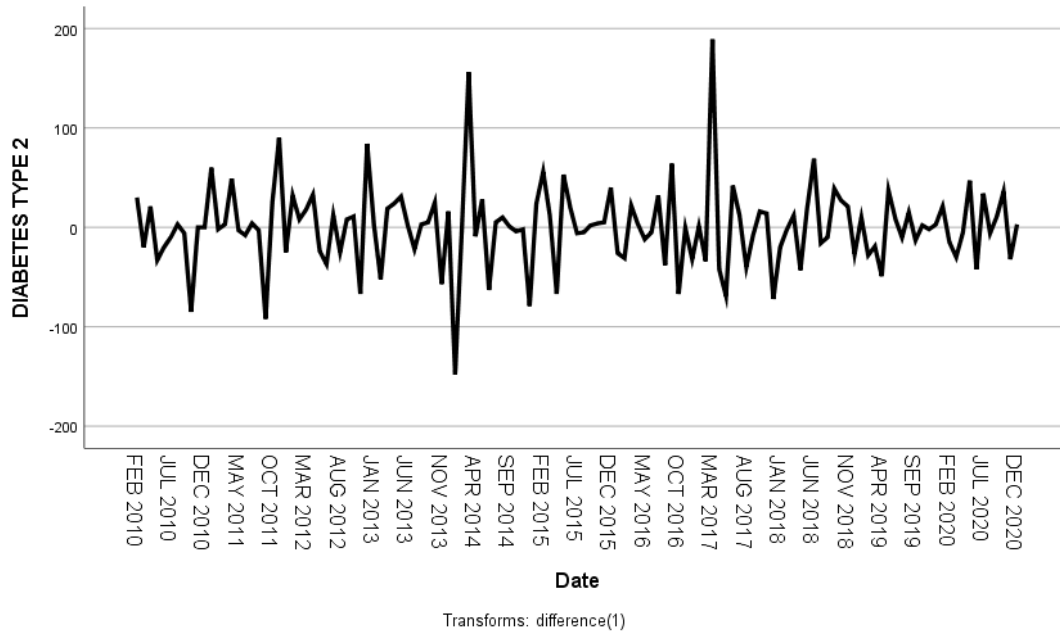


Fig. 11. Time series plot of Diabetes patients after the first differencing

Graphical Representation of Autocorrelations: From the ACF plots below (Fig. 15), the value of its function is less than one. The parameter of any given function is less than one; it means that there is stationarity.

The plot shows a slight negative spike at lag 13 and there is no other spikes showing that it is over differenced. Further the plot decays to zero on the both sides of the mean.

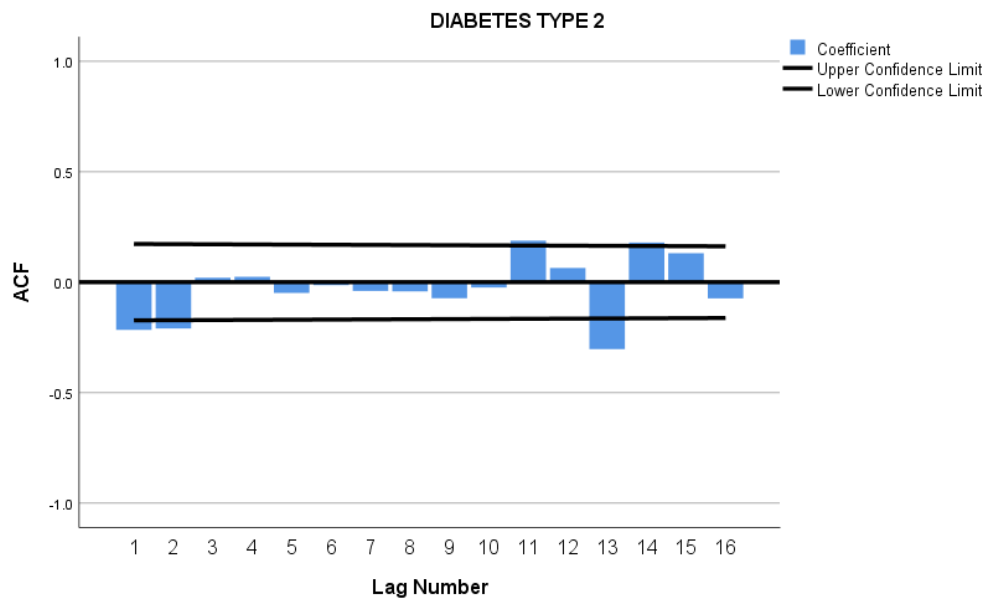


Fig. 12. Sample ACFplot of monthly Diabetes type2 Data after first differencing.

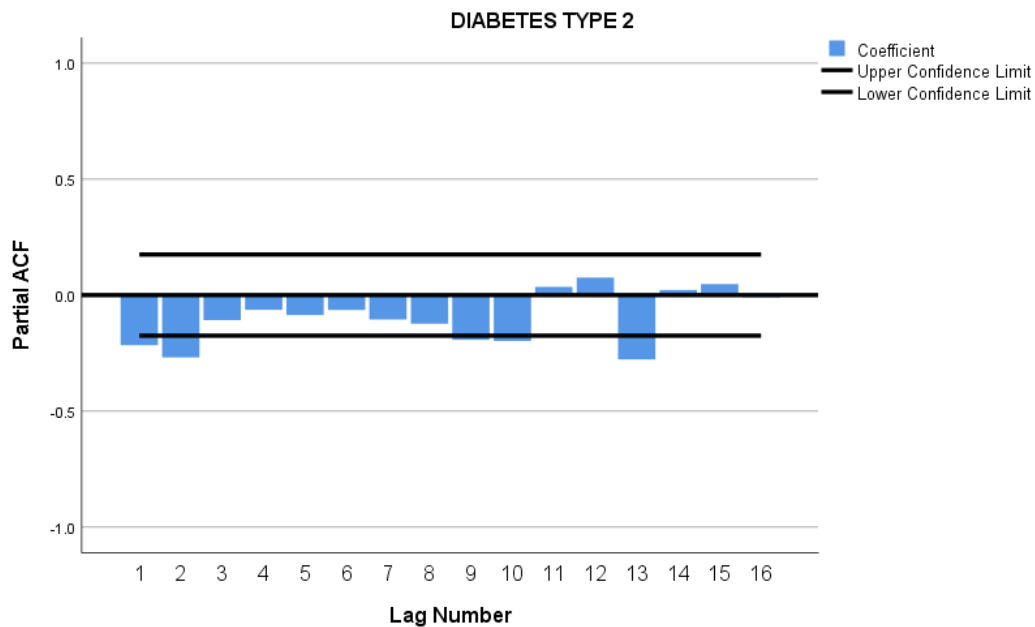


Fig. 13. Sample PACF plot of monthly Diabetes type 2 Data after first differencing

Parameter Estimation: Four models are chosen and tested to find a model with good fit. These models include ARIMA (1,1,1), ARIMA (1,1,0), ARIMA (0,1,1), and ARIMA (0,1,0).

Table 11. ARIMA model parameters

				Estimate	SE	t	Sig.
DIABETE S TYPE 2- Model_1	DIABETES TYPE 2	No Transformation	Constant	.290	.207	1.401	.164
			AR Lag 1	.553	.084	6.568	.000
			Difference	1			
			MA Lag 1	.999	.644	1.551	.123

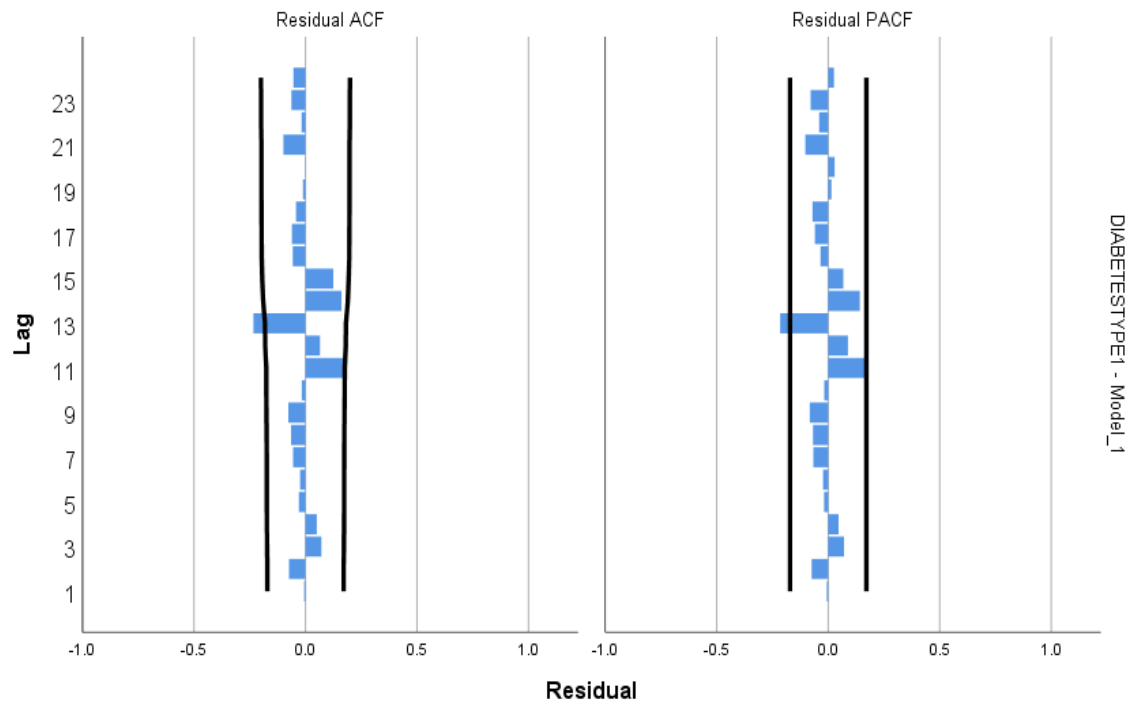


Fig. 14. Residual plots of ACF and PACF after estimating ARIMA (1,1,1)

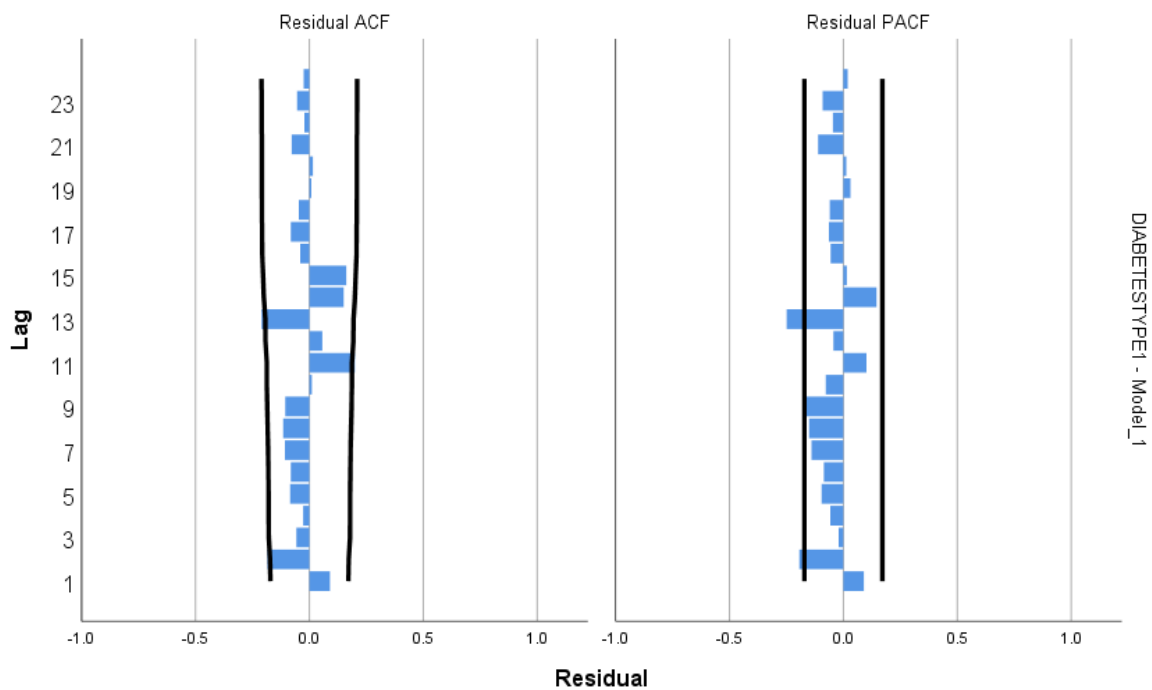


Fig. 15. Residual plots of ACF and PACF after estimating ARIMA (0,1,1)

Based on the Ljung-Box Q statistics, the models with significant p-values are selected and shown in Table 18 below. The resultant BIC providing in Table 19 shows that the most appropriate model to fit diabetes type 2 patients' data is ARIMA (1, 1, 1).

Table 12. Model Fit of ARIMA (0,1,1)

Fit Statistic	Mean	SE	Minimum	Maximum	Percentile						
					5	10	25	50	75	90	95
Stationary R-squared	.092	.	.092	.092	.092	.092	.092	.092	.092	.092	.092
R-squared	.197	.	.197	.197	.197	.197	.197	.197	.197	.197	.197
RMSE	39.755	.	39.755	39.755	39.755	39.755	39.755	39.755	39.755	39.755	39.755
MAPE	31.745	.	31.745	31.745	31.745	31.745	31.745	31.745	31.745	31.745	31.745
MaxAPE	1159.469	.	1159.469	1159.469	1159.469	1159.469	1159.469	1159.469	1159.469	1159.469	1159.469
MAE	27.847	.	27.847	27.847	27.847	27.847	27.847	27.847	27.847	27.847	27.847
MaxAE	171.586	.	171.586	171.586	171.586	171.586	171.586	171.586	171.586	171.586	171.586
Normalized BIC	7.440	.	7.440	7.440	7.440	7.440	7.440	7.440	7.440	7.440	7.440

Table 13. Model Parameters of ARIMA (0,1,1)

				Estimate	SE	t	Sig.
DIABETES	DIABETES	No Transformation	Constant	.132	2.019	.065	.948
TYPE 2-Model_1	TYPE 2		Difference	1			
			MA Lag 1	.422	.080	5.269	.000

Table 14. Model fit ARIMA(1,1,0)

Fit Statistic	Mean	SE	Minimum	Maximum	Percentile						
					5	10	25	50	75	90	95
Stationary R-squared	.047	.	.047	.047	.047	.047	.047	.047	.047	.047	.047
R-squared	.157	.	.157	.157	.157	.157	.157	.157	.157	.157	.157
RMSE	40.741	.	40.741	40.741	40.741	40.741	40.741	40.741	40.741	40.741	40.741
MAPE	31.491	.	31.491	31.491	31.491	31.491	31.491	31.491	31.491	31.491	31.491
MaxAPE	1160.286	.	1160.286	1160.286	1160.286	1160.286	1160.286	1160.286	1160.286	1160.286	1160.286
MAE	27.415	.	27.415	27.415	27.415	27.415	27.415	27.415	27.415	27.415	27.415
MaxAE	181.515	.	181.515	181.515	181.515	181.515	181.515	181.515	181.515	181.515	181.515
Normalized BIC	7.489	.	7.489	7.489	7.489	7.489	7.489	7.489	7.489	7.489	7.489

Source:spss

Table 15. ARIMA model parameters of ARIMA(1,1,0)

				Estimate	SE	t	Sig.
DIABETES	DIABETES No		Constant	.147	2.934	.050	.960
TYPE 2-	TYPE 2	No	AR Lag 1	-.215	.086	-2.500	.014
Model_1	Transformation		Difference	1			

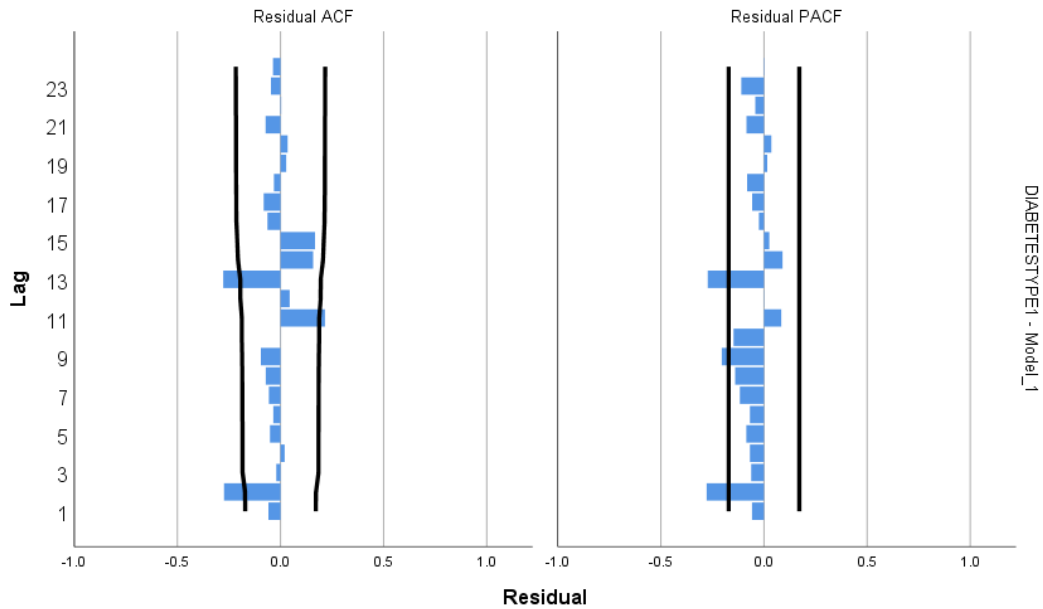


Fig. 16. Residual plots of ACF and PACF after estimating ARIMA (1,1,0)

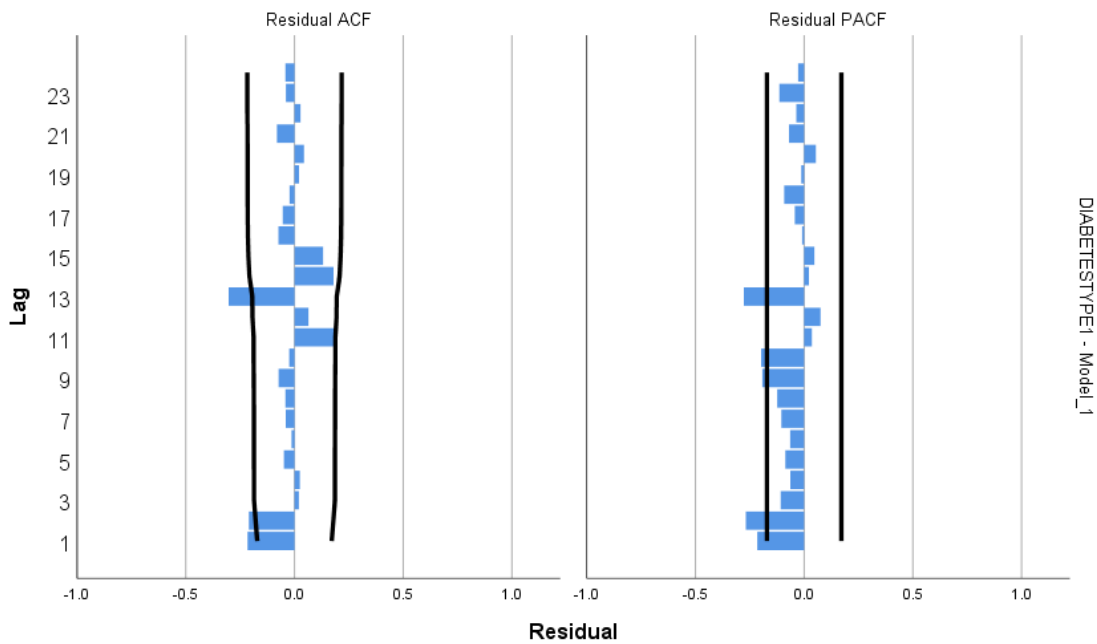


Fig. 17. Residual plots of ACF and PACF after estimating ARIMA (0,1,0)

Table 16. Model Fit of ARIMA(0,1,0)

Fit Statistic	Mean	SE	Minimum	Maximum	Percentile						
					5	10	25	50	75	90	95
Stationary R-squared	.000	.	.000	.000	.000	.000	.000	.000	.000	.000	.000
R-squared	.116	.	.116	.116	.116	.116	.116	.116	.116	.116	.116
RMSE	41.562	.	41.562	41.562	41.562	41.562	41.562	41.562	41.562	41.562	41.562
MAPE	32.512	.	32.512	32.512	32.512	32.512	32.512	32.512	32.512	32.512	32.512
MaxAPE	1152.38	.	1152.385	1152.385	1152.385	1152.385	1152.385	1152.385	1152.385	1152.385	1152.385
MAE	27.970	.	27.970	27.970	27.970	27.970	27.970	27.970	27.970	27.970	27.970
MaxAE	188.809	.	188.809	188.809	188.809	188.809	188.809	188.809	188.809	188.809	188.809
Normalized BIC	7.492	.	7.492	7.492	7.492	7.492	7.492	7.492	7.492	7.492	7.492

Table 17. Model Parameters of ARIMA(0,1,0)

				Estimate	SE	t	Sig.
DIABETES TYPE 2-Model_1	DIABETES TYPE 2	No Transformation	Constant	.191	3.631	.053	.958
			Difference	1			

Table 18. Competitive models for the monthly diabetes type 2 patients based on the Ljung-Box Q statistics test

Models	Q-statistics	P-value
ARIMA(1,1,1)	24.311	.083
ARIMA(0,1,1)	34.899	.006
ARIMA(1,1,0)	41.814	.001
ARIMA(0,1,0)	41.991	.001

Table 19. Competitive models for the monthly diabetes type 2 patients

Fit statistic	ARIMA(1,1,1)	ARIMA(0,1,1)	ARIMA(1,1,0)	ARIMA(0,1,0)
Stationary R-square	.215	.092	.047	.000
R-square	.306	.197	.157	.116
RMSE	37.114	39.755	40.741	41.562
MAPE	30.204	31.745	31.491	32.512
MaxAPE	1107.781	1159.469	1160.286	1152.385
MAE	26.553	27.847	27.415	27.970
Normalized BIC	7.340	7.440	7.489	7.492

3.8 Diagnostic checking and parameter estimation

Form Fig. 14 the sample ACF and PACF of the model shows that the autocorrelations of the residual are all close to zero which mean they are uncorrelated, hence the residual assume mean of zero and constant variance. Finally, the p-value (0.83) for the Ljung-Box statistic clearly exceeds 5% for all lag orders. Thus, the selected model ARIMA (1, 1, 1) satisfies all the model assumptions.

3.8.1 Forecasting using ARIMA (1, 1, 1)

The forecast values with 95 percent forecast limit of the ARIMA (1, 1, 1) of model for monthly diabetes type 2 Patients are shown in Fig. 18 with lower and upper limit and its actual forecast.

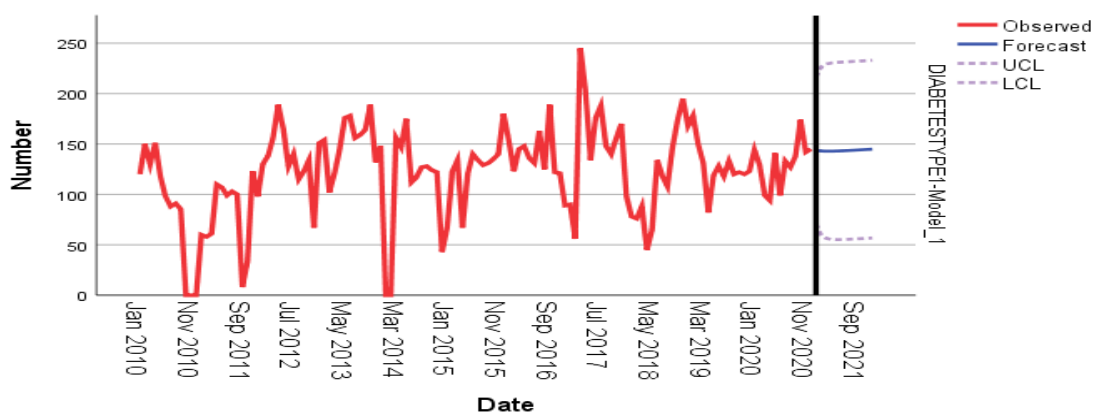


Fig. 18. Forecast chat of ARIMA(1,1,1)

4 Conclusion

In this research, Diabetes type 1 and type 2 patients were studied using the ARIMA modeling and forecasting method. Box-Jenkins was used to study the monthly Diabetes type 1 and type 2 patients records in Jos University Teaching Hospital for the period of January 2010 to December 2020. The purpose of this research work is to forecast the monthly diabetes type 1 and type 2 cases for the coming period of January 2020 to

December 2025. The results also show the forecast for next five years. List of tentative ARIMA models are developed based on the Ljung-Box Q statistic test, BIC and after testing the significant of the estimated parameters, ARIMA(3, 1, 1) model was seen best fit for forecasting the Diabetes type 1 patient and ARIMA(1,1,1) was best fit for forecasting diabetes type 2 patients. The study came out with two better fit models for predicting the prevalence of diabetes type 1 and type 2 recorded in Jos Plateau state, these models are ARIMA(3,1,1) for type 1 and ARIMA(1,1,1) for type 2. The forecast of these models shows that both diabetes type 1 and type 2 increase gradually over time, this will give researchers and health workers information on the prevalence of diabetes type 1 and type 2. Therefore, the proposed model will help to plan appropriately and allocate resources for emergency.

Competing Interests

Authors have declared that no competing interests exist.

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