Platelet lymphocyte ratio a diagnostic challenge in pancreatic head masses

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Abstract

Background: Masses in head of pancreas may arise from Carcinoma head of pancreas or Chronic pancreatitis. There is a need for differentiating the type of pancreatic head masses with utmost priority, since it can convert a radical surgery to conventional procedure. Recent studies suggest that Platelet lymphocyte ratio (PLR) can be used as a marker for differentiating between these masses. We evaluated the role of PLR and other tumour markers with combinations in differentiating the masses in head of the pancreas. Methods: A total of Sixty patients with mass in the head of pancreas with size 2cm or greater were included in the study, with 30 patients in the Pancreatic ductal adenocarcinoma (PDA) group and 30 patients in the Chronic Pancreatitis group. Patients with evidence of acute pancreatitis, liver cirrhosis, cholangitis and obstructive jaundice, having Lewis antigen A/B blood types, with absent tissue/cytological diagnosis were excluded. Results: Haematological parameters like platelet count, neutrophil count, lymphocyte count were not statistically significant between the two groups, whereas, Mean CA 19-9 levels, (PLR) Platelet lymphocyte ratio levels and (NLR) Neutrophil lymphocyte ratio in (PDA) Pancreatic ductal adenocarcinoma were higher than Chronic pancreatitis group. (P<0.05) In all these patients, biomarkers such as CA 19-9, Platelet lymphocyte ratio (PLR), Neutrophil lymphocyte ratio (NLR) and different combinations of these tumour markers were calculated, and their sensitivity, specificity,(PPV) Positive predictive value, (NPV) Negative predictive value and accuracy in differentiating malignant from benign pancreatic masses were established. The diagnostic accuracy of the tumour markers and their combinations are in the following order PLR> PLR+CA 19-9> CA 19-9+ NLR > NLR. Conclusion: We found that PLR is as good as tumour marker CA 19-9 both individually as well as with other tumour marker combinations in the diagnosis of pathology of pancreatic head masses. Cost effectiveness, availability, simplicity and better diagnostic accuracy made PLR better tumour marker in the current scenario.

Keywords: Platelet lymphocyte ratio, Carcinoma head of pancreas, Chronic pancreatitis, Ca19-9

Introduction

Masses in the Head of the pancreas always pose a diagnostic dilemma to differentiate the nature, as they may arise from Carcinoma head of pancreas or Chronic pancreatitis. PDA (pancreatic ductal adenocarcinoma) develops in the head of the pancreas causing majority of cancer-related deaths owing to its aggressiveness. However, early diagnosis of such condition improves the survival rate. Some malignancies can also arise from the head of the pancreas secondary to chronic pancreatitis; it is wise to keep surveillance in patients with benign aetiology. Many Tumour markers like CA 19-9, tissue inhibitor of metalloproteinase-1 (TIMP-1), intercellular adhesion molecule 1 (ICAM 1) are available to diagnose the malignancy of pancreas [1-3]. Among them, CA 19-9, a sialylated Lewis blood group antigen has shown its consistency with well accepted sensitivity and specificity which varies with various cut-off values - (<37ku/l<81% & 90%), (100 ku/l-98% &68%), (1000ku/l-99.8% & 41%) [4]. However, CA 19-9 has got false negative results in conditions like Lewis blood type a, b and false positive results in obstructive jaundice and large tea consumers [5-7]. Goonetilleke et al. stated that CA 19-9 should be used in the contemporary algorithm for the diagnosis of pancreatic malignancies [8]. The role of hematological parameters like Platelet lymphocyte ratio and Neutrophil lymphocyte ratio as tumor markers are subject of interest for diagnosis and prognostication [9, 10]. PDA causes...
thrombocytosis and leukocytopenia. Tumour produces cytokines (IL-6) which stimulate thrombocytosis as a part of paraneoplastic thrombocytosis. High platelets prevent tumour cells destruction from NK-cells and also helps in Tumour cells extravasation and angiogenesis. Clinically thrombocytosis may precede the diagnosis of PDA by months/years. Platelet lymphocyte ratio is also helpful in diagnosing the patients developing malignancy from an inflammatory setting, better than CA 19-9 [11].

R K. Miglani in a study stated that, PLR can be used for both diagnostic and prognostic purposes [12]. Smith et al [13] found that Pre-operative PLR is an independent predictor of survival on multivariate analysis. To this purpose, we evaluated the role of PLR and other tumour markers with combinations in differentiating the masses in head of the pancreas.

Materials & Methods

This prospective study was conducted during May-2013 to April-2016 at Narayana medical college & Hospital, Nellore.

Sample Size- A total of 60 patients with 30 patients in Pancreatic ductal adenocarcinoma group and 30 patients in Chronic pancreatitis group were included.

Inclusion Criteria- Sixty patients who were admitted with mass in the head of the pancreas with size 2cm or greater were included.

Exclusion Criteria- Patients with evidence of acute pancreatitis, liver cirrhosis, cholangitis and obstructive jaundice, having Lewis antigen A/B, with absent tissue/cytological diagnosis were excluded.

Study Method- Demographic, personal and family histories were recorded from all the patients. Various risk factors for developing PDA and Chronic pancreatitis were gathered from the history such as family history with at least two affected first-degree relatives, environmental risk factors like Tobacco Chewing, Smoking and Alcoholism and medical risk factors like Pancreatitis, Diabetes (>5yea), and Obesity. Patients underwent CBP with PLR & NLR, LFT, CA 19-9, USG Abdomen MDCT with Triphasic CT and CT-Angiography for assessment of resectability. PDA patients were grouped into resectable, borderline resectable and advanced disease.

Only 5/30 patients were found out to be resectable and underwent Whipple procedure with R0 Resection. 3/30 patients were considered borderline resectable and within these patients, 2/30 patients had R1 Resection, and 1/30 had R0 Resection. 22/30 patients who had locally and distant metastasis, the diagnosis was made by intraoperative tissue biopsy while contemplating triple bypass and CT guided FNAC of metastasis.

In chronic pancreatitis, the majority of the patients without dilated main pancreatic duct underwent FREY procedure, and 4/30 patients with dilated MPD underwent BEGERS procedure. 6/30 patients who had adhesions due to recurrent pancreatitis underwent Pylorus-Preserving Pancreaticoduodenectomy.

Statistical Methods- Continuous variables were presented as the mean, median and standard deviation. Categorical variables were described in frequencies / percentiles. We used MaxStat statistical software for data processing and analysis. Comparison of the continuous variables of the study group was made using students’ t test and Mann-Whitney "U" test. Fisher's exact test was used to analyse absolute values.

Sensitivity, specificity, positive predictive value, negative predictive value and accuracy of each tumour markers and their combinations were calculated. Statistical significance was defined when P value < 0.05.
Table-1: Demographic, clinical and haematological parameter comparison between the two disease groups.

| Clinical data        | PDA          | Chronic pancreatitis | P value |
|----------------------|--------------|----------------------|---------|
| 10-20 Yrs            | 0            | 1(3.33%)             | <0.05   |
| 21-30 Yrs            | 1(3.33%)     | 4(13.32%)            |         |
| 31-40 Yrs            | 2(6.67%)     | 8(26.64%)            |         |
| 41-50 Yrs            | 7(23.31%)    | 8(26.64%)            |         |
| 51-60 Yrs            | 8(26.64%)    | 6(19.98%)            |         |
| >60 Yrs              | 12(39.96%)   | 3(9.99%)             |         |
| Male: Female Ratio   | 21:9 (2.3:1) | 24:6 (4:1)           | >0.05   |
| Family history       | 4(12%)       | 2(6%)                | >0.05   |
| Smoking              | 12(40%)      | 5(15%)               | <0.05   |
| Tobacco chewing      | 2(6%)        | 4(12%)               | >0.05   |
| Alcoholism           | 7(21%)       | 27(91%)              | >0.05   |
| Obesity              | 13(39%)      | 10(30%)              | <0.05   |
| Type-2 DM (>5years)  | 9(27%)       | 11(33%)              | >0.05   |
| Mean platelet count  | 2.93±1.15    | 2.42±1.21            | >0.05   |
| Mean lymphocyte count| 1601±330     | 2028±407             | >0.05   |
| Mean Neutrophil count| 2587±225    | 2370±384             | >0.05   |
| CA19-9               | 589±499      | 102±101              | <0.05   |
| PLR                  | 227±128      | 121±97               | >0.05   |
| NLR                  | 3.65±1.39    | 1.25±1.17            | <0.05   |

Table-2: Sensitivity, specificity, positive predictive value, negative predictive value and accuracy of each tumour markers and their combinations

| Tumour markers and their combinations | Sensitivity | Specificity | PPV | NPV | Accuracy |
|---------------------------------------|-------------|-------------|-----|-----|----------|
| CA 19-9                               |             |             |     |     |          |
| >37u/l                                | 91          | 50          | 69  | 92  | 68       |
| >100u/l                               | 80          | 69          | 96  | 79  | 69       |
| >1000u/l                              | 74          | 97          | 99  | 76  | 73       |
| Platelet Lymphocyte ratio             |             |             |     |     |          |
| <150                                  | 78          | 86          | 78  | 77  | 65       |
| >150                                  | 71          | 90          | 80  | 64  | 70       |
| Neutrophil Lymphocyte Ratio           |             |             |     |     |          |
| <2.5                                  | 67          | 70          | 66  | 70  | 56       |
| >2.5                                  | 60          | 72          | 69  | 64  | 66       |
| Combinations of tumour markers        |             |             |     |     |          |
| CA19-9+PLR                           | 72          | 88          | 74  | 67  | 68       |
| CA 19-9+NLR                          | 60          | 72          | 68  | 63  | 56       |
| NLR>2.5                               | 42          | 61          | 55  | 49  | 43       |

Discussion

Pancreatic head masses always impose a diagnostic conundrum to the clinician. The obvious quagmire is to differentiate malignant and benign head masses.

Tumour markers are quite essential in early diagnosis, assessing the severity, prognosis and surveillance of the disease [14]. (ASCO) American Society of Clinical Oncology in its 2006 update concluded that CA 19-9 alone with its sensitivity and specificity is not adequate for diagnosis of pancreatic cancer [15]. National Academy of Clinical Biochemistry (NACB) suggested that CA19-9 should be combined with imaging modalities to increase the efficacy and strongly recommended studies for newer tumour markers to identify high-risk individuals, and with better sensitivity and specificity.

Tumour markers like CA19-9, PLR & NLR and combinations were taken into this predicament, and PLR
showed superlative efficacy. Leukocytopenia is due to release of inflammatory cytokines (IL-1, IL-6 & TNF-alpha) which augment the cancer cells to metastasize by affecting the CRI (cancer related inflammation) cascade relating Lymphocytopenia with tumour aggressiveness/metastasis [9-11]

Smith et al., found that high pre-op PLR levels in suspected periampullary carcinoma with extensive tumour invasiveness and concluded that 1/5th of laparoscopic surgeries could be avoidable if the combination of these markers is considered [16]. In cases of resected PDA, pre-op PLR was found to be more important tumour marker than other haematological parameters. The pre-op PLR found to be significant in multivariate analysis (p<0.001) along with tumour size (p<0.001) and lymph node ratio (p<0.013) [16].

Rk Miglani et al. showed that- PLR can be used for both diagnostic and prognostic values and can differentiate malignant masses from benign aetiology. When used in combination the accuracy had increased [12].

In another study conducted by Bhatti et al., NLR retained significance in resected PDA than PLR [17]. In a study, the borderline resectable PDA patients were grouped based on pre-op PLR & NLR for survival analysis and found that the NLR>3 and PLR>225 were independent prognostic factors in Borderline resectable PDA patients.

In our study, we have taken consideration of tumour markers like CA 19-9, PLR, NLR and their combinations. Many studies had proved CA19-9 as a diagnostic and prognostic tumour marker, but the level of acceptance by the clinician varies as the accuracy varied by various cutoff values. PLR(>150) was found to be a single independent prognostic factor on multivariate analysis with (p<0.005) and the accuracy of the various combinations are PLR> PLR+CA 19-9> CA 19-9+PLR> NLR> CA 19-9+NLR> PLR> was supportive in case of malignancy arising from chronic pancreatitis in 3 patients, however, is insignificant (p>0.005).PLR alone was found to be simple cost-effective and accurate diagnostic tumour marker than other tumour markers and combinations which yielded poor results.

Conclusion

We found that PLR is as good as tumour CA 19-9 both individually as well as with other tumour marker combinations in the diagnosis of pathology of pancreatic head masses. Cost effectiveness, availability, simplicity and better diagnostic accuracy made PLR better tumour marker in the current scenario.

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