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# Early Virological Response of the First Line Combination Therapy (Pegylated Interferon ? -2a and Ribavirin) in Iraqi Chronic Hepatitis C Patients and Their Psychological Adverse Effects Dr. Vian Ahmed Wasta Ismael<sup>1</sup> Hawler Medical University Received: 11 February 2012 Accepted: 29 February 2012 Published: 15 March 2012

#### 9 Abstract

This study was designed to assess short-term therapeutic effectiveness and psychological 10 adverse effects of combination of pegylated interferon ?-2a and ribavirin in Iraqi chronic 11 hepatitis C patients. For this purpose fifty newly diagnosed chronic hepatitis C patients 12 divided into three groups A, B and C, treated with equal doses of pegylated interferon ?-2a 13 (180 ?g/week) and different doses of ribavirin (1200, 1000 and 800 mg/day respectively) and 14 followed up for 12 weeks of starting treatment (prospective groups). Twenty healthy subjects 15 were selected to be a normal group for the purpose of comparison. The results at week 12 (the 16 time of achieving EVR) showed 100 17

18

#### 19 Index terms—

#### <sup>20</sup> 1 I. Introduction

he hepatitis C virus (HCV) is a major public health problem and a leading cause of chronic liver disease (Williams
R, 2006). Hepatitis C is the principal cause of death from liver disease and the leading indication for liver
transplantation in the United States (Kim W, 2002).

Approximately 20-30% of patients with chronic HCV infection progress to end stage liver disease within 20 years and a small percentage develop hepatocellular carcinoma (Walker R and Edwards O, 2007). HCV infection is now the leading worldwide indication for liver transplantation (Ryan K and ??ay C, 2004, Walker R andEdwards O, 2007)

<sup>27</sup> O, 2007). First-line treatment for HCV includes pegylated interferon plus ribavirin. The dosing regimen varies with the 28 specific product and the duration of therapy varies with the product and HCV genotypes (Wells B et al, 2006). 29 Interferon (IFN) was first described in 1957 as an antiviral compound in chick embryo cells (Greenwood D 30 et al, 2007). Its efficacy for treatment of HCV first was recognized when Hoofnagle et al (1986) published a 31 preliminary findings when HCV was known as non-A, non-B hepatitis (Sangik O and Afdhal N, 2006). The 32 United States Food and Drug Administration (FDA) approved alpha interferon monotherapy for the treatment 33 34 of chronic HCV infection in 1992. Ribavirin was approved for use as an adjunct to interferon therapy of hepatitis 35 C in 1998. Pegylated forms of interferon in combination with ribavirin were approved in the United States in 36 2001 (Hoofnagle J, 2009). There are two licensed pegylated interferons, peg interferon ?-2b (Peg-Intron, Schering Plough Corp.), with a 12-kd linear polyethylene glycol (PEG) covalently linked to the standard interferon ?-2b 37 molecule, and peg interferon ?-2a (Pegasys, Hoffmann-La Roche) with a 40-kd branched PEG covalently linked 38 to the standard interferon ?-2a molecule (Zeuzem S et al, 2003). The doses of these two forms of pegylated 39 interferons (Peg IFNs) differ (Ghany M et al, 2009). Peg IFN ?-2b is dosed according to body weight (1.5 ?g/kg 40 once weekly), while the larger Peg IFN ?-2a is given in a fixed dose of 180 ?g once weekly (Cornberg M et al, 41 2002). 42

Peg IFN ?-2b may also be dosed at 1.0 ?g/kg once patients become negative for HCV-RNA without major
declines in sustained virological response (SVR) rates (McHutchison J et al, 2009, Manns M et al, 2011).

Pegylation of interferons increases the persistence of the interferon in the blood, extend half life, better toleration and much importantly produces much superior virological response (Greenwood D et al, 2007).

The two licensed peginterferons have been shown in head-to-head comparison to be equivalent in efficacy and to have similar safety profiles (McHutchison Ribavirin is a synthetic nucleoside in which ribose is linked to a triazole derivative. Like other nucleoside analogues it has to be activated intracellularly by phosphorylation (Greenwood D et al, 2007). The precise mode of action has proved elusive, though there are several theories including the possibility that it causes lethal mutations in viral nucleotides (Greenwood D et al, 2007). Ribavirin has limited utility as monotherapy and should be administered twice daily with food when used in combination with ?-interferons (DiPiro J et al, 2005).

Meta-analyses and systematic reviews confirm that a combination of PegIFN with RBV is effective in treating patients with chronic hepatitis C (CHC), leading to high levels of SVR (Strader D et al, 2004). In general, the combination of RBV with ?-interferons is associated with numerous adverse events to multiple organ systems, and these should be discussed with patients prior to initiation of therapy (DiPiro J et al, 2005).

This study was designed to measure early virological response (ERV) in HCV infected patients receiving first line combination therapy (Peg interferon ?-2a and ribavirin) and to evaluate response according to different doses of ribavirin but fixed dose of peg interferon ?-2a. Also this study was conducted to monitor appearance of psychological adverse effects to guide the patients and provide necessary instructions.

# <sup>62</sup> 2 II. Subjects and Methods

## <sup>63</sup> **3** a) Patients

This study was conducted during the period from the 15th March 2012 till 1st October 2012, which was carried
out in Gastro-enterology center at General teaching hospital in Sulaimania city. Fifty six patients (30 males and
26 females) with an age of 18-70 years were divided into three groups according to viral genotypes (36 patients

<sup>67</sup> infected with genotype 1 and 20 patients infected with genotype 4), dose of the ribavirin and body weight of the <sup>68</sup> patients. Throughout the study period, six patients were lost (4 males and 2 females) and only fifty patients

(26 males and 24 females, in whom 32 patients were infected with genotype 1 and 18 patients with genotype 4)

<sup>70</sup> followed up. Four of the six patients stopped taking the drug (poor adherence), one female died (car accident),

<sup>71</sup> and the other female withdraw the drug after one month of treatment because of severe dehydration, arthralgia,

myalgia, head ache, nausea and vomiting. All patients were recieving combination of 180 ?g/week of Peg IFN
 ?-2a s.c. injection (Pegasys <sup>®</sup> by Roche pharmaceutical company, Switzerland) and different doses of ribavirin

?-2a s.c. injection (Pegasys ® by Roche pharmaceutical company,
capsules (Rebetol ® by Schering pharmaceutical company, USA).

For monitoring of hematological and other common adverse effects from combination therapy that may necessitate dose adjustment or even withdrawal of the drugs, the patients were examined weekly for the first month then monthly for the other 2 month.

Ethical authorization and permission were submitted from each of college of pharmacy, directory of health
 and gastroenterology center in Sulaimania city. Informed concern had been taken from patients studied.

80 The previously diagnosed patients were recruited into the following prospective groups:

Group A : This group included fourteen patients infected with HCV genotype one, 10 males and 4 females ranging 22-65 years (mean  $\pm$  SD, 45.4  $\pm$  12.15), with body weights more than 75 kg, taking combination of PegIFN ?-2a 180 ?g once weekly as subcutaneous injection and RBV capsule 1200 mg per day (three 200 mg capsules after breakfast and three 200 mg capsules after dinner).

Group B : Included eighteen patients infected with HCV genotype one, 9 males and 9 females ranging 21-67 years (44.94  $\pm$  14.7), with body weights equal or less than 75 kg, taking combination of Peg IFN ?-2a 180 ?g once weekly as subcutaneous injection and RBV capsule 1000 mg per day (three 200 mg capsules after breakfast and two 200 mg capsules after dinner).

Group C : Included eighteen patients infected with HCV genotype four, 7 males and 11 females ranging 18-65
years (43.78 ± 13.28), taking combination of Peg IFN ?-2a 180 ?g once weekly as subcutaneous injection and
RBV capsule 800 mg per day (two 200 mg capsules after breakfast and two 200 mg capsules after dinner).

# 92 4 b) Healthy Subjects

Twenty healthy individuals were involved as a control group, including 8 males and 12 females ranging 19-69 years  $(39.1 \pm 13.4)$ .

# <sup>95</sup> 5 c) Inclusion criteria

Patients confirmed to have HCV infection, genotypes 1 and 4. ? Patients between 18-70 years old of both
 genders.

- 98 ? Treatment naïve patients.
- 99 ? Patients willing to be treated and to adhere to treatment requirement.

# <sup>100</sup> 6 d) Exclusion criteria

101 ? Patients with HIV or HBV co-infection.

Patients with solid organ transplantation (heart, lung, liver, and kidney) ? Patients with decompensatedliver disease.

Patients allergic to any one of the components of combination therapy. ? Difficult to follow up patients (alcoholics, patients who travel frequently). ? Breast feeding and pregnancy or patients unwilling to comply with adequate contraception.

107 ? Patients with severe psychiatric disorder.

108 ? Patients with severe immunosuppression.

109 ? Patients with heart failure or significant coronary or CVD. ? Patients with untreated thyroid disease.

110 ? Patients with unknown HCV genotype (refused to do viral genotyping).

e) Sample collection and preparation 4 ml of venous blood was collected from each patient. The blood was

drawn by venipuncture under basal condition using tourniquet with vacationer system, then centrifuged at 3000 rpm to seperate plasma and stored in ACD or EDTA tube and freezed at -20? (within 4 hrs of collection) and

<sup>113</sup> rpm to seperate plasma and stored in ACD or EDTA tube and freezed at -20? (within 4 hrs of collection) and <sup>114</sup> analyzed within 2 week. The assessments were done twice for each patient, once before starting treatment and

second time three months after starting treatment with combination therapy of peg IFN ?-2a and RBV.

# <sup>116</sup> 7 f) Sample processing and extraction

Purification of viral nucleic acid from cell was carried out using genomic DNA extraction method in which nucleic 117 acids of the virus are lysed quickly and efficiently using lyses buffer which is a highly concentrated solution of 118 chaotropic salt. When combined with ethanol, the buffer creates optimum conditions for nucleic acid binding to 119 the glass fiber matrix of the column tube. Contaminants such as salts, metabolites and soluble macromolecular 120 cellular component are removed in the washing step. Nucleic acid is eluted in RNAase-free water and is then 121 ready for use in subsequent reactions including real time RT-PCR and other enzymatic reactions (Bioneer Inc., 122 2009). g) Amplification of viral nucleic acid Amplification of viral nucleic acid (RNA for HCV) was carried 123 out by real time RT-PCR procedure using EXICYCLER ® (BIONEER/ South Korea). RNA templates are 124 first reverse-transcribed to generate complementary cDNA strands followed by a DNA polymerase-mediated 125 cDNA amplification. DNA detection simultaneous to amplification is preferentially achieved by the use of 126 target sequence-specific oligonucleotides linked to two different molecules, a fluorescent reporter molecule and 127 a quenching molecule. These probes bind the target cDNA between the two PCR primers and are degraded 128 or released by the DNA polymerase during DNA synthesis. In case of degradation the reporter and quencher 129 molecules are released and separated, which results in the emission of an increased fluorescence signal from the 130 reporter. The fluorescence signal, intensified during each round of amplification, is proportional to the amount 131 of RNA in the starting sample (Mauss S et al, 2012). 132

## <sup>133</sup> 8 h) Psychological evaluations

For evaluation of psychological conditions of HCV infected patients before treatment and three months after 134 starting treatment with 180 ?g/week of s.c. peg IFN ?-2a and different doses of RBV, each patient were interviewed 135 and filled a used questionnaire. The questionnaire was prepared by a psychologist Dr. Rebwar H. Gharib at 136 2008 for his research (Gharib R, 2008) using DSM-IV scoring system (American Psychiatric Association, 1994). 137 According to the questionnaire, patients who presented with at least 5 of depressive symptoms during the same 138 two-week period or more, at least one of which is either depressed mood or loss of interest, is considered to have 139 major depression. Patients who presented with at least 3 of depressive symptoms during the same two-week 140 period or more, at least one of which is either depressed mood or loss of interest, is considered to have minor 141 depression. Insomnia, suicidal idea and suicidal attempt were also considered separately. 142

# <sup>143</sup> 9 i) Statistical analyses

All data are represented as mean ± standard error of means (SEM). Statistical analysis were carried out using
paired sample T-test to compare treatment groups, focusing on changes from pre-treatment values and after three
months of starting treatment of each group. Statistical analyses were carried out using SPSS 16.

## 147 **10 III. Results**

#### <sup>148</sup> 11 a) Effects of combination therapy on viral load

Table (1) and figure (1) show a significant reduction in viral load (amount of HCV-RNA in serum) three months after starting treatment compared to viral load before starting treatment. Nearly in all patients (47 patients), the viral load became no detectable in serum, but in only two patients (one 38 years old female, and one 36 years old male, both in group C) the virus was still detectable but comparing to pre-treatment amount there was more than two log reduction. Only one 63 years old female in group B was resistant to treatment and viral load increased by 2 times the pre-treatment value three months after starting treatment. In those patients whose serum viral RNA became no detectable, viral load reduced to  $0.0 \pm 0.0$  IU/ml compared to pretreatment values

#### 13 C) PSYCHOLOGICAL CONDITION

of  $19427000 \pm 527847$  IU/ml (very high),  $2374000 \pm 331629$  IU/ml (high),  $914420 \pm 31540$  IU/ml (moderate) and  $258500 \pm 20169$  IU/ml (low) with percent reduction of 100%.

#### 158 12 b) Other parameters

In all three groups (as showed in tables 2 and 3), there were significant reductions (p ? 0.05) in the levels of each of hemoglobin (Hb), white blood cell count (WBC), absolute neutrophil count (ANC), platelet count (PLT), random plasma glucose level, Alanine

# <sup>162</sup> 13 c) Psychological condition

In figure 2 for group A patients (n=14); before starting treatment, two patients (one male and one female) 163 were complaining from isolated insomnia (insomnia alone, without any other psychological symptoms), which 164 represents 14.3% of all group A patients, who both became suffering from major depression later three months 165 after starting treatment. And one female patient was complaining from minor depression before treatment 166 167 representing 7.1% of all the group's patients who also became suffering from major depression three months 168 after starting treatment. The remaining 78.6% were psychologically normal patients. After 3 months of starting treatment with s.c. Peg IFN ?-2a 180 ?g/week and RBV 1200 mg/day, three cases (21.4%) of minor depression, 169 170 four cases (28.6%) of major depression one of whom also had suicidal idea, three cases (21.4%) of isolated insomnia 171 were reported. Among those who reported major depression, one patient was male (who also had a suicidal idea) and the other three were females, all three patients with minor depression were male, and those with isolated 172 insomnia were two males and one female. Overall, three months after starting treatment, only four patients 173 (28.6%) were not complaining from psychiatric symptoms (three males and one female), while the remaining ten 174 patients (71.4%) were complaining from psychiatric symptoms as individualized above. 175

In figure 3 for group B patients (n=18); before starting treatment, six patients (one male and five females) 176 177 were complaining from minor depression which represents 33.3% of all group B patients, one female of whom 178 had suicidal idea. Later, three months after starting treatment, all of these six patients became suffering from major depression, two females of these became having suicidal idea without suicidal attempt. And only one 179 180 female patient representing 5.6% was complaining from isolated insomnia, who became majorly depressed and died later after three months of starting treatment as a result of suicidal attempt. The remaining 61.1% were 181 psychologically normal patients. After 3 months of starting treatment with s.c. Peg IFN ?-2a 180 ?g/week and 182 RBV 1000 mg/day, seven cases (38.8%) of minor depression, five cases (27.8%) of major depression three of 183 whom had suicidal idea, three cases (16.7%) of isolated insomnia were reported. Unfortunately, 5 days after 184 my interview with patients, one 43 years old female who had isolated insomnia alone before starting treatment, 185 committed suicide by jumping out of a building, after three days of staying at hospital, she passed away. One 186 187 point of note is that, this female had experienced major depression and frequent suicidal idea compared to the 188 pre-treatment state, before committing suicide. Among those who reported major depression, two patients were male and the other three were females, two males and five females were with minor depression, and those with 189 isolated insomnia were two males and one female. Overall, three months after starting treatment, only three 190 patients (16.7%) were not complaining from psychiatric symptoms, while the remaining fifteen patients (83.3%) 191 were complaining from psychiatric symptoms as individualized above. 192

In figure 4 for group C patients (n=18); before starting treatment, two female patients were complaining from 193 minor depression which represented 11.1% of all group C patients, one of them was also complaining from suicidal 194 idea. One female (5.6%) was complaining from major depression before treatment who interestingly became 195 minor depressed three months after starting treatment. No one was complaining from isolated insomnia before 196 197 treatment. The remaining 83.3% were psychologically normal patients. After 3 months of starting treatment with s.c. Peg IFN ?-2a 180 ?g/week and RBV 800 mg/day, four cases (22.2%) of minor depression, five cases 198 (27.8%) of major depression one of whom had suicidal idea and tried to commit suicide, four cases (22.2%) 199 of isolated insomnia were reported. Unfortunately, three days before my interview with patients, one 54 years 200 old female who already had minor depression and suicidal idea even before starting treatment, tried to commit 201 suicide by burning herself, but she was lucky and rescued by her son who prevented her from doing such a thing. 202 One point of note is that, at the time of interview, three days before trying to commit suicide, this female had 203 experienced major depression and more frequent suicidal idea compared to the pre-treatment state. Among those 204 who reported major depression, one patient was male and the remaining four were females, two males and two 205 females were with minor depression, and those with isolated insomnia were two males and two females. 206

Overall, three months after starting treatment, only five patients (27.8%), two males and three females, were not complaining from psychiatric symptoms, while the remaining thirteen patients (72.2%) were complaining from psychiatric symptoms as individualized above.

In figure 5 for control group (n=20); Two patients (one male and one female) were complaining from minor depression which represents 10% of all control group individuals, one female with major depression and suicidal idea which represents (5%), and seven patients (two males and five females) representing 35% were complaining from isolated insomnia, the remaining 50% were psychologically normal individuals.

Finally in figure 6 we may say that, among all three groups of patients (A, B, and C), percent of patients experienced major depression and percent of those who were psychologically normal three months after starting treatment, were greater in group A (28.6% for each) compared to group's B and C patients. For minor depression, the percent of patients experiencing it, was greater in group B patients (38.8%) compared to other groups, and isolated insomnia was more frequent among group C patients than group A and B patients, and percent of patients experienced major depression were the same in both groups B and C (27.8%). One point of note, percent of individuals experiencing isolated insomnia was greater (35%) among control group than other three patient groups after three months of treatment.

# <sup>222</sup> 14 IV. Discussion

Hepatitis C virus infection is still a global and the possible new approaches for conquering the health care 223 challenges. The standard of care (SOC) therapy for patients with chronic hepatitis C virus infection has been 224 the use of both peg interferon and ribavirin (Ghany M et al, 2011). These drugs are administered for either 48 225 weeks (HCV genotypes 1, 4, 5, and 6) or for 24 weeks (HCV genotypes 2 and 3), inducing sustained virological 226 response rates of 40%-50% in those with genotype 1 and of 80% or more in those with genotypes The therapy 227 of hepatitis C began almost 26 years ago with a small trial of recombinant human interferon alfa (Hoofnagle J 228 229 et al, 1986). The rationale for using interferon was its broad antiviral effects and the suspicion that it might be 230 active against the stillundiscovered agent of non-A non-B hepatitis. Not until the discovery of the HCV, at 1989 (Feitelson M, 2003), were the effects of interferon understood. Nevertheless, interferon was approved for use for 231 232 hepatitis C treatment in the United States in 1992 (Hoofnagle J, 2009).

The second important advance in hepatitis C therapy came with the use of ribavirin. Ribavirin was approved for use as an adjunct to interferon therapy of hepatitis C in 1998. A third advance in therapy of hepatitis C came soon thereafter, with the introduction of pegylated forms of interferon that allowed for onceweekly (rather than thrice-weekly) injections. Peg nterferon was approved in the United States in 2001 (Hoofnagle J, 2009). The treatment paradigm for HCV has changed with the recent FDA approval of two first generation protease inhibitors, telaprevir, and boceprevir for genotype 1 infected individuals. Nonetheless, ribavirin and pegylated interferon remain integral components of treatment (Ghany M et al, 2011).

In our community, combination of peginterferon and ribavirin is still the first choice because of high cost of triple therapy and difficulties in providing these drugs on continuous bases. Despite that, majority of our patients cannot afford such a large amount of money for providing the drugs themselves continuously. So in order to determine advantage and effectiveness of this first line combination therapy (by measuring early virological response which is a main predictor for sustained virological response in majority of patients) in HCV infected patients in our communuity, its psychological effects after three months of treatment, present study has been conducted.

The results of present study are somewhat conflicting and out of line with the results of other studies because patients treated in clinical trials, represent a highly selected population not necessarily representing general HCV infected population (Ferenci P et al, 2005). Therefore, it is not clear if the reported efficacy and safety of peg interferon ? and ribavirin regimen would be validated in routine clinical practice.

Response to standard treatment with peg interferon ?-2a and ribavirin in patients with chronic hepatitis caused by HCV, including genotypes 1 and 4, has been widely studied and documented in numerous populations (Fried M et al, 2002a, Hadziyannis S et al, 2004). Approximately 80% of patients who have genotype 1 and virtually all patients who have genotypes 2 and 3 achieve an early virological response (Davis G, 2002, Fried M et al, 2002a, Lindsay K, 2002, Shiffman M et al, 2007a).

Early virologic response (EVR) was defined as the ? 2 log 10 reduction in HCV RNA in serum 12 weeks after starting treatment. In case of total absence of HCV RNA in serum 12 weeks after starting treatment, a complete early virologic response (cEVR), which is a more promising predictor of sustained virological response (SVR) than EVR, is obtained (Mauss S et al, 2012). Over all, in this study among all participants, 94% achieved cEVR (100%, 94.4% and 88.9% for groups A, B and C respectively) 4% achieved ? 2 log 10 reduction in HCV RNA, i.e., EVR (11.1% of group C patients) and 2% null responder (5.6% of group B patient) who discontinued drugs after 12 weeks of starting treatment after confirming increased viral load, patient's instruction and acceptance.

In a retrospective analysis done by Gheorghe L et al (2005) In one of a phase III trials of peg interferon ?-2a 263 and ribavirin, by week 12 of therapy, EVR was achieved by 86% of patients (Fried M et al, 2002a) which is near 264 to present study results. These results suggest that patients who have EVR who remain PCR positive at 12 265 weeks (not complete absence of HCV RNA) should have PCR testing repeated after 24 weeks before making any 266 decision about discontinuing therapy. Achievement of EVR can provide a goal to motivate patient adherence 267 during the first months of therapy, and early testing provides the opportunity to reassess the need for continued 268 269 treatment. Consequently, when an EVR is absent, discontinuation of therapy should be considered because the 270 likelihood of sustained response is negligible, but the decision must be made on an individual patient basis. If 271 uncertainty exists, retesting should be considered before stopping therapy (McHutchison J and Fried M, 2003, 272 Manns M, 2004), which was the case with the female patient in group B in this study, when repeated viral load testing after one week of last PCR showed the same increase in HCV RNA compared to baseline level. 273

In a study done by Ascione A et al (2010) in Italy including both genotypes 1 and 4, EVR was obtained in 85% of all patients. The majority of patients obtained a complete EVR, while the number of those who obtained a partial EVR was only 8.8%. The results of Ascione A et al's study are nearly the same as that in present study. In the registration trials of peg interferon ?-2a plus ribavirin, 10% to 14% of patients had to discontinue therapy due to an adverse event (Manns M et al, 2001, Fried M, 2002b), compared to 2% discontinuation in present study. Laboratory abnormalities are the most common reasons for dose reduction. Among these, neutropenia (absolute neutrophil count [ANC] of 1500 mm<sup>3</sup>) was a frequent laboratory abnormality, occurring in 18% to 20% in the two large phase III clinical trials where the dose was reduced 50% for an ANC of 750 mm<sup>3</sup> and permanently discontinued for an ANC of ? 500mm<sup>3</sup> (Manns M et al, 2001, Fried M et al, 2002a).

Severe neutropenia, ANC ?500 mm<sup>3</sup>, occurred in 4% of subjects. None of these lab abnormalities were reported in present study participants. Actually these abnormalities happened in participants but not so severe to necessitate dose reduction but if study participants were followed up further (i.e., more than three months) these severe effects that call for dose modification may appear. Interferon causes anemia via bone marrow suppression and ribavirin causes anemia via hemolysis ??De Franceschi L et al, 2000).

For psychological presentation, among all 50 patients 28% presented with major depression 10% of whom had suicidal idea and 4% committed suicide. 28% presented with minor depression, 20% with isolated insomnia and the remaining 24% were psychologically normal HCV infected patients.

Major depression was more common in group a patients (28.6%) than group's B and C patients (27.8% for 291 each group), minor depression was more common in group B patients (38.8%) than group A patients (21.4%) 292 and group C patients (22.2%). Isolated insomnia was more common among group C patients (22.20%) as well 293 294 as control group (35%) compared to group a patients (21.4%) and group B patients (16.7%). These results 295 demonstrate that appearance of any type of depression or psychological symptoms is more related to individual's 296 susceptibility for developing symptoms and surrounding environment, because as it obvious, all three groups of patients treated with equal doses of peg interferon ?-2a but the percent of patients showed psychological 297 presentations differ, and its known that it is interferon that induce psychological abnormalities not ribavirin, 298 that's why psychological presentations in any one of the groups is not related to the variations in ribavirin dosing 299 regimen. Only one interesting female patient in group C, who experienced major depression before starting 300 treatment, became minorly depressed at 12 th week of treatment. The pre-treatment depression in this patient 301 was because of knowing that she is infected with HCV and misunderstood by some specialists that it is not 302 curable and should be isolated from family and friends so as not to infect others. But later when we explained 303 the disease course, routes of transmission and precautions that needed to be made, she felt better but still minor 304 depressed after three months this one may be the effect of interferon on neurotransmitters. 305

Manns M et al (2006) demonstrated that depression in HCV-infected individuals occurs in up to 60% (Hilsabeck 306 R and Malek A, 2004, Zacks S et al, 2006). During HCV treatment with interferon based regimens the prevalence 307 of depression has been reported to be between 10%-40% depending on the screening method used (Zacks S et al, 308 2006). Recently, data from the Virahep-C study, a prospective analysis of depression during HCV genotype 1 309 treatment with peg interferon and ribavirin, demonstrated that low social support was independently associated 310 with pretreatment and on-treatment development of depression (Evon D et al, 2009). At any time during the 311 Virahep-C study 20% had depressive symptoms whereas 18% of non-responders compared to 10% of responders 312 had depression 6 months after the end of treatment, a difference that may be explained by the failure of a 313 challenging 48-week treatment course. This explains that occurrence of depression is more common during first 314 three months of treatment, the time when the patients are still unaware of their response rate to treatment. In 315 Virahep-C study 21% of patients developed depression during the first 12 weeks, a result which is nearly the 316 same as present study findings. 317

A recent study of 1010 HCV infected patients demonstrated that suicide risk was higher in males and in 318 patients under the age of 45 (Kristiansen M et al, 2011), this finding is in contrary with that in present study, 319 because the two patients who had suicidal idea and committed suicide (one died and the other one prohibited) 320 were both females and the other three patients who had suicidal idea but did not try to commit it were 2 females 321 and 1 male and all were above 40 years old. Different conclusions were reported in a study of almost 400 HCV-322 infected patients with genotype 1 on peg interferon and ribavirin, where just 3.5% reported suicidal ideation and 323 none attempted suicide (Evon D et al, 2009), this may be due to extensive supervision and guidance of patients 324 by psychiatrists and family involvement strategy but neither is done during present study period. 325

Elsewhere in the literatures, many HCV treatment studies have reported various depression and suicidal ideation rates, making the application of these results to clinical practice impractical. This inconsistency in reporting may reflect the utilization of different depression screening methods, physician and patient biases in diagnosing and reporting symptoms, and variable treatment protocols followed. It is important to acknowledge the effect of interferon on the thyroid and the potential development of depressive-like symptoms related to thyroid dysfunction, mimicking, or even masking depressive symptoms related to interferon use (Papafragkakis H et al, 2012).

In a cross-sectional study of 43 patients who had chronic hepatitis C and not receiving interferon ?. Kraus M 333 et al (2001) found several factors that correlated significantly with depression. Higher rates of depression were 334 observed in older patients ( $^{250}$  years; P = 0.024), in patients who were aware of their hepatitis diagnosis for more 335 than 5 years (P = 0.003), and in patients who were informed that they were not eligible for interferon? therapy 336 (P = 0.001). Furthermore, a lower incidence of depression was noted in patients who had been diagnosed with 337 HCV recently (1-6 months; P = 0.003). If such evaluations were done in present study in HCV infected patients 338 before starting treatment, at least some if not all explanations made by Kraus et al may be applicable especially 339 age more than 50 years and newly diagnosed ones. 340

Finally we should not exclude the major role of patient care and need of clinical pharmacist interventions in 341 improving patient's adherence and response to standard therapy. Interim results of a prospective, randomized, 342 controlled multicenter study indicate that active intervention with patient education, aggressive side effect 343 management, and expanded supportive nursing intervention with cognitive behavioral therapy by way of telephone 344 calls for patients infected with HCV who are treated with Peg-IFN and RBV therapy is feasible, can decrease the 345 dropout rate in the first 12 weeks of therapy, and is associated with significant improvements in patient quality 346 of life at early time points in treatment ??Sarrazin C et al, 2010a). In present study some if not all of mentioned 347 above strategies followed, so this may be the main cause of high rate of EVR especially cEVR and low rate of 348 unwanted effects of treatment on organ functions. 349

# 350 15 V. Conclusions

? First line combination therapy with Pegylated interferon ?-2a and ribavirin is highly effective in early eradication
of hepatitis C virus (98% EVR) in Iraqi chronic hepatitis C patients and can be used relatively safely . ? Lower
doses of ribavirin lowers percent of patients who achieve complete EVR. NO. of patients

# <sup>354</sup> 16 Psychiatric symptoms





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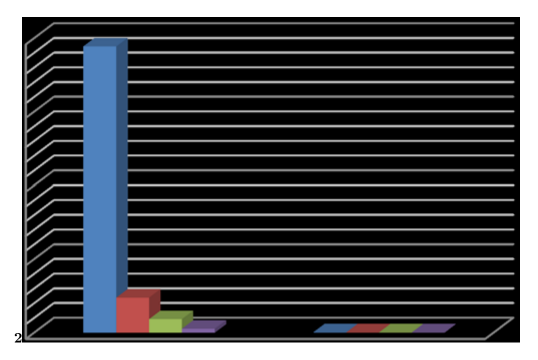


Figure 2: Figure 2 :

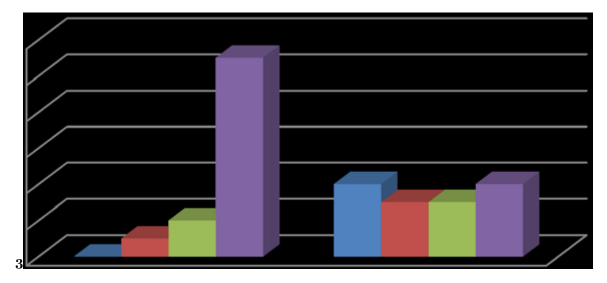


Figure 3: Figure 3 :

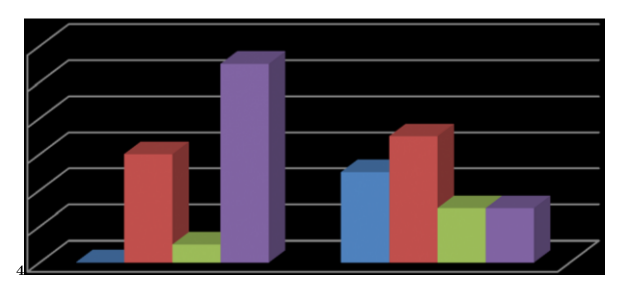
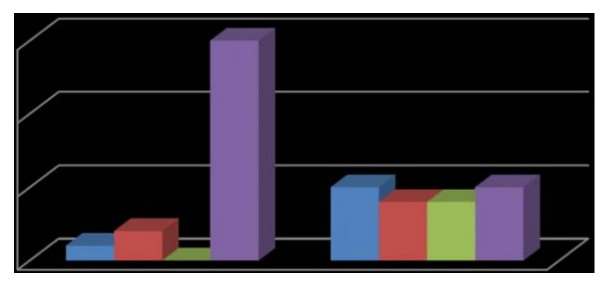


Figure 4: Figure 4 :





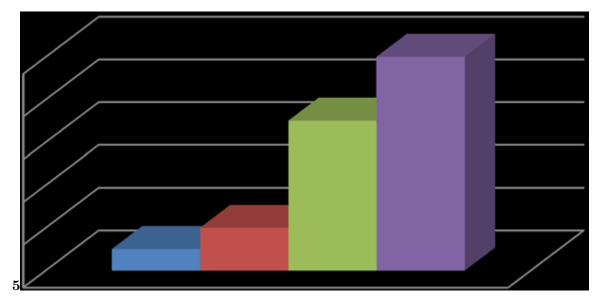


Figure 6: Figure 5 :

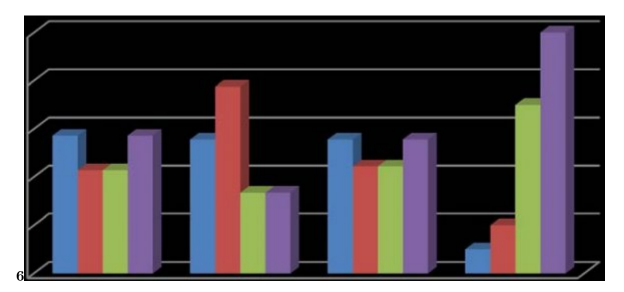


Figure 7: Figure 6 :

? Patients with thalasemia, cytopenia, or severe aminotransferase (ALT), Alkaline phosphatase (ALP), anemia. serum albumin and body weight three months after

? Patients with renal failure. starting treatment compared to pre-treatment values. Neither dos**r**edu**ntiph**armacological

interventions were required, since hematological reductions were not severe.

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Figure 8:

ear 2012 Y Medical Research Volume XII Issue XI Version I Global Journal of in Romania that consisted of 174 HCV infected patients, therapy was stopped in patients who do not achieve 2 log reductions in viral load 12 weeks after starting treatment, the same strategy that followed in present study. Early virological response (EVR) in a clinical trial at Beth Israel Medical Center NY (Johnson T et al, 2004) was 63% in genotype1 patients treated with peg

Figure 9:

Figure 10:

50.00%				
40.00%				
30.00%				
20.00%				
10.00%				
	Group A	Group B	Group C	Control
Major depression	Group A 28.60%	Group B 27.80%	Group C 27.80%	$\begin{array}{c} \text{Control} \\ 5\% \end{array}$
Major depression Minor depression	-	*	-	0 0 0 -
	28.60%	27.80%	27.80%	5%

Figure 11. Democrat	of notionta	Darroological	armontoma
Figure 11: Percent	or patients	Psycological	symptoms

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- [Table 1 : Viral load (amount of HCV RNA in serum in IU/ml) of HCV infected patients, genotypes one and four, before treatmed Table 1 : Viral load (amount of HCV RNA in serum in IU/ml) of HCV infected patients, genotypes one and
- four, before treatment and 3 months after starting treatment with PegIFN ?-2a 180 ?g/week and RBV 1200,
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