**Original Research Article**

**Fibrinogen levels helps in early detection of abnormal pregnancies**

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**ABSTRACT**

**Background:** Haemostatic failure as an end result of various complications of pregnancy is an important cause of maternal mortality in India. The main aim of this study is to detect the levels of fibrinogen in abnormal pregnancy i.e. Pregnancy induce hypertension (PIH), Intra uterine fetal death (IUFD), Missed abortion, Abruptio placenta.

**Methods:** Study was conducted on 150 in patients joined in Obstetrics ward private hospital, Andhra Pradesh, 50 are control subjects and 100 are study patients, Out of hundred (100) cases, 40 are PIH, 25 are IUFD, 25 are Missed abortion, and 10 are Abruptio placenta. Estimated for fibrinogen, D-Dimer, total proteins, Albumin.

**Results:** The fibrinogen levels in present study decreased significantly. PIH (Control mean 442.0, S.D ±43.38, Test mean 296.0, S.D ±48.03, p<0.001). IUFD (Control mean 442.0, S.D±43.38, Test mean 262.4, S.D±20.06, p<0.001). Missed abortion (Control mean 442.0, S.D ±43.38, Test mean 250.80, S.D±26.13, p<0.001). Abruptio placentae (Control mean 442.0, S.D±43.38, Test mean 210.5, S.D±87.38, P<0.001). D-dimer levels are estimated semi quantitatively and the levels were found to be increased. Total proteins and albumin are decreased in all the cases, but significantly in PIH (T.P-Control mean6.25, S.D±0.65, Test mean 5.25, S.D±1.57, p<0.001, Albumin- Control mean 2.79, S.D±0.34, Test mean 2.23, S.D±0.59, p<0.001).

**Conclusions:** The estimation of plasma fibrinogen is helpful not only in the early diagnosis of haemostatic failure but also to guide replacement therapy during the fibrinopenic state.

**Keywords:** Abruptio placenta, IUFD, Missed abortion, PIH

**INTRODUCTION**

Haemostatic failure as an end result of various complications of pregnancy is an important cause of maternal mortality in India. Incidence of maternal mortality in India due to complications during pregnancy and child birth is 17.5% (Out of 3, 58,000 globally 63,000 deaths occur in India.)

Understanding of abnormalities of coagulation parameters and their significance in various obstetric emergencies mandates thorough knowledge of not only normal physiology of blood clotting mechanism but also physiological changes in pregnancy and pathophysiology of such obstetric conditions. In following sections, we shall discuss etiopathogenesis and clinical significance of coagulation abnormalities in missed abortion, abruptio placentae, pregnancy induced hypertension (PIH) and intra uterine fetal demise (IUFD).

**Physiology of blood coagulation**

Coagulation factors in blood are present in inactive state and an appropriate stimulus like endothelial damage or tissue damage prompts activation of coagulation cascade which limits loss of blood by formation of blood clot. Simultaneously another system of anticoagulants work in tandem with clotting system to limit unwanted extension of this clot to normal sites. Delicate balance of both these systems is necessary to maintain homeostasis.
The loose aggregation of platelets in temporary plug is bound together and converted into the definitive clot by fibrin. The clotting mechanism responsible for formation of fibrin involves a cascade of reactions in which inactive enzymes are activated, and the activated enzymes in turn activate other inactive enzymes.

The fundamental reaction in the clotting of blood is conversion of soluble plasma protein fibrinogen into insoluble fibrin.

**Physiological changes during pregnancy**

Pregnancy is a physiological state, where changes occur virtually in every organ system and blood is not an exception. Coagulation system changes create a hypercoagulable state. Main reason for this state is increase in plasma levels of fibrinogen. Normal levels of fibrinogen are 200-400 mg/dl, where as in pregnancy fibrinogen levels reach up to 600 mg/dl. Also there is increase in activities of factors II, VII, VIII, IX, X, X13. Anticoagulants like protein C, protein S and anti-thrombin III are present in decreased levels. As a result of this, there is increase in fibrinogen and other clotting factors and depressed fibrinolytic activity, a hypercoagulable state is created. Accelerated intra vascular coagulation serves to maintain utero-placental interface. These changes in coagulation system minimize blood loss by effective haemostatic mechanisms after separation of placenta.

**Disseminated intravascular coagulation (DIC)**

DIC is an acute or chronic thrombo hemorrhagic disorder occurring as a secondary complication of variety of diseases. It is characterized by activation of coagulation sequence that leads to formation of microthrombi throughout the micro circulation of the body. As a consequence of the thrombotic diathesis, there is consumption of platelets, fibrin and other coagulation factors. Thus this disorder is also called consumption coagulopathy. Secondly, there is activation of fibrinolytic mechanisms to remove the clots from microcirculation. Etiology of DIC is wide and includes various obstetric and non-obstetric conditions show in Table 1.

The consequences of DIC are, first there is wide spread deposition of fibrin within the micro circulation leading to tissue hypoxia and ischemia. Fragmentation of red blood cells squeezing from these vessels can lead to hemolysis (micro angiopathic hemolytic anemia). Second, as result of uncontrolled clot formation, platelets and coagulation parameters are consumed at rate greater than the rate of their production. This leads to hemorrhagic diathesis. In addition fibrinolysis leads to formation of fibrin degradation products, which inhibit platelet aggregation and fibrin polymerization. All these lead to haemostatic failure in DIC.

Here we shall discuss four obstetric conditions which are associated with abnormalities on coagulation parameters.

- Intra uterine fetal death (IUFD)
- Abruptio Placenta
- Pregnancy Induced Hypertension (PIH)
- Missed Abortion.

**Intra uterine fetal death (IUFD)**

Involuntary loss of pregnancy at any period of gestation is fetal demise. Fetal death as defined by WHO in 1950 and revised by the working group formed by the American academy of pediatrics and American college of Obstetricians and Gynecologists in 1998, is “death prior to complete expulsion or extraction from its mother of a products of human conception, irrespective of the duration of pregnancy and which is not an induced termination of pregnancy”. The death is indicated by the fact that after such expulsion or extraction, the fetus does not show any evidence of life such as heart beats, definite movements of voluntary muscles, umbilical cord pulsations.

Clinically IUFD is suspected when mother reports loss of fetal movements. Clinical signs like gradual regression of fundal height, loss or diminished uterine tone and inability to feel fetal movements and inability to hear fetal heart sound support the diagnosis of IUFD. Egg shell cracking of fetal head is a late sign. The diagnosis of IUFD can be confirmed by ultrasound Doppler by noting the absence of fetal heart activity.

It has been discussed above that if fetus is retained for more than 4 weeks (as in 20% of cases) there is possibility of defibrination from silent disseminated intravascular coagulation (DIC). This is due to gradual absorption of thromboplastin, liberated from dead placenta and decidua, into the maternal circulation. Although IUFD might lead to DIC, frequently the underlying cause for IUFD such as HELLP syndrome, preeclampsia or placental abruption themselves might be associated with DIC.

Incidence of coagulopathy in IUFD is determined by 2 factors cause of IUFD and duration of retained dead fetus.

**Abruptio placenta**

Separation of the placenta from its implantation site before the birth of fetus is called Abruptio placenta. This entity has been variously termed as accidental hemorrhage and ablation placenta in the literature.

Its incidence is 1.5% in all pregnancies and it falls to 0.3% in full term pregnancies. Incidence of abruptio placenta in India has been found to be 0.2 to 2%. Retroplacental clot is detected post-partum in 4.5% of cases.
The initial event in abruptio placenta is bleeding into decidua basalis. The retro-placental hematoma/ hemorrhage formed separate the placenta from the maternal vascular system causing impairment in fetal oxygenation and nutrition.

It has been suggested that, first, a retroplacental clot forms and may build up sufficient pressure to cause abruptio placentae. In full blown abruptio placentae, uterine size may be larger than term uterus and contains a large retroplacental clot which might have tracked beyond the confines of placental margin. Blood may track down and bleeding per vaginum can be a presenting symptom. Then it is called revealed abruptio placentae. Bleeding may be intrauterine and no blood appears at vulva, called concealed abruptio placentae. More commonly a mixed variety is seen with both the features. In concealed abruptio placentae blood clot may even burst into amniotic sac (port wine colored amniotic fluid). The blood may dissect into myometrium towards serosa resulting in full blown picture of Couvelaire uterus (uteroplacental apoplexy). It may show characteristic echhymosis on the serosa. Bleeding may occur between the layers of broad ligament. Peritoneal cavity contains blood stained fluid and in rare cases, uterus may actually rupture into peritoneal cavity leading to massive intra peritoneal bleed.

The resulting laceration in the decidual layers also allows free communication from the intra decidual space of the hematoma into the maternal circulation of the placenta. There now follows a process in which tissue substances including thromboplastin from decidual layers enter directly into maternal circulation leading to coagulopathy.

As evident from above discussion, abruptio placentae can be associated with abnormalities in coagulation cascade ranging from mild changes to fulminant DIC. The grade of abrupton and magnitude of abnormalities in lab parameters correlate to a large extent and hence DIC can be seen in cases of severe abruptio placentae.

**Pregnancy induced hypertension (PIH)**

There is no antecedent history or any documentation that the woman had hypertension prior to pregnancy. The hypertension develops as a direct result of gravid state. It is divided into three clinical types:

- **Pre-eclampsia**
- **Eclampsia**
- **Gestational Hypertension**

Incidence in hospital practice varies widely from 5-15%. The incidence in primigravidae is about 10% and in multigravidae 5%.

In pre-eclampsia an imbalance of different components of prostaglandins- relative or absolute deficiency of vasodilator prostaglandin (PGI2) synthesized in vascular endothelium and increased synthesis of thromboxane (TXA2), a potent vasoconstrictor in platelets.

**Coagulation:** Evidence of disseminated intravascular coagulopathy DIC affecting wide spread organs, due to release of trophoblastin into the circulation. It may arise from the blood platelets as in Shwartzman reaction or from release of trophoblastic fragments into the uterine circulation. There is reduction of platelets, Fibrinogen, antithrombin III and plasminogen levels in the blood. Microthrombi affect the arterioles of all the vital organs apart from the placenta. Degree of thrombocytopenia reflexes the severity of pathology. Fibrinectin and thrombin levels are elevated.

**Missed abortion**

**Definition of abortion**

Abortions is the termination of pregnancy before the period of viability which is considered to occur at 28th week. The limit of viability is brought down to either 20th week or fetus weighing 500gm.

**Incidence:** The incidence of miscarriage is about 10%. 75% abortion occurs before 16th week and of these about 75% occurs before the 8th week of pregnancy.

**Blood coagulation disorders:** If fetus is retained for more than 4 weeks there is possibility of defibrination from silent disseminated intravascular coagulopathy (D.I.C). It is due to gradual absorption of trophoblastin, liberated from dead placenta and decidua, into the maternal circulation. 3. Psychological upset 4. During labor-Uterine inertia, retained placenta and postpartum haemorrhage.

The main aim of this study is to detect levels of fibrinogen levels in the abnormal pregnancy. Which is main blood component in clot formation, other parameters like D-dimer, total protein and albumin are also estimated.

**METHODS**

In the present study 150 cases admitted in obstetric ward for delivery were studied (control 50 and cases 100) with following parameters, Plasma fibrinogen, d-dimer, Total protein, Albumin. The mean age group of patients is 25 years, during the period of 2014-2015. Out of hundred pregnancies forty 40 are PIH, 25 are Missed abortion, 25 are IUD, 10 are Abruptio Placenta. Fifty (50) healthy control of normal pregnancy who are admitted for delivery of similar time. Statistics analysis is done for p value.

Plasma fibrinogen is estimated by Tulip fibroquant method.
Procedure for fibrinogen calibrator curve

1. The thrombin reagent vial reconstituted exactly with one ml of DW wait for 15 min. Now it is ready to use for the fibrinogen test.
2. The Fibrinogen Calibrator vial reconstituted with exactly one ml of DW, Wait for 15 min. This is the Fibrinogen Calibrator stock solution.
3. 1:5, 1:10, and 1:20 etc., dilutions of fibrinogen calibrator solution are prepared (eg: 0.05ml Std+ 0.95ml Buffer for 1:20, 0.1ml+ 0.9ml of std and Buffer resp. for 1:10 dilution etc) and added as in the table.
4. Pipette into labelled test tubes A, B, C, D, and E for standards and do.

Take 0.2ml of Fibrinogen Calibrator dilution into test tube from each std tube and add 0.1ml of reconstituted thrombin reagent and simultaneously start stopwatch.
Stop the stopwatch at the first appearance of the fibrin web, as the gel clot beings to form and record the time in seconds.
The time against the fibrinogen concentration is plotted on the graph paper Fig: 1 provided by the kit itself and points are connected which gave a straight line.

Procedure for testing the sample

Plasma 100μl
Owren’s Buffer 900μl
Take 200μl from the above solution and incubate at 37°C for one minute and add 100μl of pre warmed (37°C) thrombin reagent and immediately start the stopwatch.
Stop the stopwatch as the gel clot being to form and record the time in seconds.

If fibrinogen content is high the clotting time will be less and if the less fibrinogen clotting time will be more.

Normal Range: 150mg-400mg/dl

D-Dimer is estimated by Tulip XL FDP D-Dimer

Semi quantitative method

- Immediately start the stopwatch. Rock the slide gently, back and forth, observing for agglutination macroscopically at three minutes.
- Agglutination in the highest plasma dilution corresponds to the approximate amount of D dimer level in ng/ml.
- To calculate D dimer level in ng/ml in the sample, use the following formula.
- D dimer level ng/ml= 200×d
  - d= highest dilution of plasma showing agglutination during the semi quantitative test of the sample.

Total protein and albumin is estimated quantitatively in fully automated analyser EM360 Transasia by using ERBA kits.

RESULTS

The fibrinogen levels in present study decreased significantly. PIH (Control mean 442.0, S.D ±43.38, Test mean 296.0, S.D ±48.03, p<0.001). IUFD (Control mean 442.0, S.D±43.38, Test mean 262.4, S.D±20.06, p<0.001). Missed abortion (Control mean 442.0, S.D ±43.38, Test mean 250.80, S.D±26.13, p<0.001). Abruptio placentae (Control mean 442.0, S.D±43.38, Test mean 210.5, S.D±87.38, P<0.001). D-dimer levels are estimated semi quantitatively and the levels were found to be increased. Total proteins and albumin are decreased in all the cases, but significantly in PIH (T.P-Control mean6.25, S.D±0.65, Test mean 5.25, S.D±1.57, p<0.001, Albumin- Control mean 2.79, S.D±0.34, Test mean 2.23, S.D±0.59, p<0.001).

<table>
<thead>
<tr>
<th>Obstetric conditions</th>
<th>Non-obstetric conditions</th>
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<tbody>
<tr>
<td>Abruptio placenta</td>
<td>Infections</td>
</tr>
<tr>
<td>Pre-eclampsia and eclampsia</td>
<td>Neoplasms</td>
</tr>
<tr>
<td>IUFD</td>
<td>Septic abortion</td>
</tr>
<tr>
<td>Missed abortion</td>
<td>Excessive blood loss</td>
</tr>
<tr>
<td>Septic abortion</td>
<td>Puerperal sepsis</td>
</tr>
<tr>
<td>Excessive blood loss</td>
<td>Amniotic fluid embolism</td>
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</table>

| Table 1: Major disorders associated with DIC. |

| Table 2: Standardization of fibrinogen levels. |

<table>
<thead>
<tr>
<th>S_A</th>
<th>S_B</th>
<th>S_C</th>
<th>S_D</th>
<th>S_E</th>
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<tr>
<td>50μL</td>
<td>100μL</td>
<td>150μL</td>
<td>200μL</td>
<td>250μL</td>
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<tr>
<td>130</td>
<td>260</td>
<td>390</td>
<td>520</td>
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<td>950</td>
<td>900</td>
<td>850</td>
<td>800</td>
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</table>
DISCUSSION

In pregnancy complicating disease like Abruptio placentae, pregnancy induced hypertension, Intra uterine death and missed abortion common terminal complication is occurrence of DIC which is responsible for maternal mortality.

After the onset of clinical manifestation, the diagnosis of DIC is simplified but the treatment becomes difficult. Thus it is important to diagnose DIC at its subclinical stage so that early therapeutic measures can be instituted. The estimation of plasma fibrinogen is helpful not only in the early diagnosis of haemostatic failure but also to guide replacement therapy during the fibrinopenic state.

In the present study we studied 100 cases of abnormal pregnancies, out of them pregnancy induce hypertension (40), Intra uterine death (25), Missed abortion (25), Abruptio placentae (10), the plasma fibrinogen levels is decreased compared to normal pregnant woman.

In Abruptio placentae (10 cases) the fibrinogen levels in our study decreased more significantly compared to other condition (plasma levels 210mg/dl, p<0.001). Our study is supported by Memon FA, Noorani KJ, Parasnis H, Raje B, Hinduja IN, Wisot AL, Barczak EM One out of ten (10%) developed DIC.

### Table 3: Biochemical parameters in PIH.

<table>
<thead>
<tr>
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<th>Mean</th>
<th>S.d</th>
<th>T-value</th>
<th>P-value</th>
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<tr>
<td>T.P</td>
<td>6.25</td>
<td>0.65</td>
<td>1.57</td>
<td>4.053</td>
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<tr>
<td>Albumin</td>
<td>2.79</td>
<td>0.34</td>
<td>0.59</td>
<td>5.530</td>
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<tr>
<td>Fibrinogen</td>
<td>442.0</td>
<td>43.38</td>
<td>48.03</td>
<td>15.127</td>
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### Table 4: Biochemical parameters in IUFD.

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<tr>
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<td>6.25</td>
<td>0.65</td>
<td>1.03</td>
<td>1.119</td>
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<tr>
<td>Albumin</td>
<td>2.79</td>
<td>0.34</td>
<td>0.639</td>
<td>0.017</td>
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<tr>
<td>Fibrinogen</td>
<td>442.0</td>
<td>43.38</td>
<td>20.06</td>
<td>19.629</td>
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### Table 5: Biochemistry parameters in missed abortion.

<table>
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<th>P-value</th>
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<td>6.25</td>
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<td>0.45</td>
<td>1.777</td>
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<tr>
<td>Albumin</td>
<td>2.79</td>
<td>0.34</td>
<td>0.55</td>
<td>0.324</td>
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<tr>
<td>Fibrinogen</td>
<td>442.0</td>
<td>43.38</td>
<td>26.13</td>
<td>20.239</td>
</tr>
</tbody>
</table>

### Table 6: Biochemistry parameters in missed abortion.

<table>
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<th>S.d</th>
<th>T-value</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>T.P</td>
<td>6.25</td>
<td>0.65</td>
<td>0.96</td>
<td>2.554</td>
</tr>
<tr>
<td>Albumin</td>
<td>2.79</td>
<td>0.34</td>
<td>0.36</td>
<td>1.817</td>
</tr>
<tr>
<td>Fibrinogen</td>
<td>442.0</td>
<td>43.38</td>
<td>87.38</td>
<td>12.687</td>
</tr>
</tbody>
</table>

In Abruptio placenta hypofibrinogenemia is due to free entering of decidual tissue thromboplastin from decidual layers to maternal circulation leading to coagulopathy. This is due to free communication from intra decidual
space of the hematoma into maternal circulation of the placenta.\textsuperscript{2,6}

Hypofibrinogenemia associated with increased levels of fibrin degraded products (d-dimer), which inhibit myometrical contractility leading to postpartum hemorrhage. In present study d-dimer levels are estimated semi quantitatively which is raised significantly (plasma level 3200ng/ml) to normal pregnancy (plasma level (130-1700ng/ml) our study supported by by Memon FA, Noorani KJ\textsuperscript{9}, Nolan TE, Smith RP, Devoe LD.\textsuperscript{22}

The levels of total protein and albumin are decreased in abruptio placentae then the normal pregnancy.\textsuperscript{6}

![Figure 1: Fibrinogen concentration to time in second.](image1)

![Figure 2: Pattern of distribution of cases and controls in the study.](image2)

![Figure 3: Comparison of total protein levels control and cases.](image3)

![Figure 4: Comparison of albumin levels in control and cases.](image4)

![Figure 5: Comparisons of fibrinogen levels in control and cases.](image5)

![Figure 6: Comparison of D-DIMER levels in control and cases.](image6)
In Intra uterine fetal death (25 cases) in our study show hypofibrinogenemia. 10 of 25 (40%) are <4 week and 15 are >4 weeks retained fetus intra uterine, (plasma level 262mg/dl, p=0.001) our study is supported by J Womens Health (Larchmt). Maslow AD et et al.27,21, Parasnis H, Raje B, Hinduja IN23, Gochberg SH.26

In Intra uterine fetal death hypofibrinogenemia is due to gradual absorption of thromboplastin, liberated from dead placenta and decidua, into the maternal circulation.2,6,14

The fibrin degraded products levels increased d-dimer levels (plasma levels 1600ng/ml) then the normal (2nd trimester levels 320-1290ng/ml) our study supported by Bonnar J, Davidson JF, Pidgeon CF, McNicol GP, and, Douglas AS.10

In missed abortion (25 cases) in our study show hypofibrinogenemia (plasma level 250mg/dl) our study is supported by Parasnis H, Raje B, Hinduja IN23, Chandran R, Adeeb N.24

The underlying pathology for hypofibrinogenemia in missed abortion is similar to that of IUF D.13

d-dimer levels (plasma levels-1600ng/ml) then the normal (2nd trimester levels 320-1290ng/ml), total protein, albumin levels are decreased.

In pregnancy induce hypertension (40 cases) in our study show hypofibrinogenemia (plasma levels 296mg/dl). 15 of 40 (37.5%) show normal levels. Our study supported by Yang M, Shen W, Chen L20, Schjetklein R, Haugen G, Wilsolffv, Parasnis H, Raje B, Hinduja IN23, Trott EL.29, Mary Part Fitzgerald, Floro C, Siegel J, MD, Enrique Hernandez.32

D-dimer levels are increased in PIH plasma levels 800ng/dl or normal (2nd trimester levels 320-1290ng/ml) study supported by Eichinger S Nolan TE, Smith RP, Devoe LD.22

Total protein (serum level 5.25mg/dl p<0.001) decreased and albumin decreased (serum level 2.23mg/dl, p<0.001) Seong WJ, Chong GO, Hong DG, Lee TH, Lee YS, Cho YL, Chun SS and Park IS, Mack HC.6,31

**CONCLUSION**

The estimation of plasma fibrinogen is helpful not only in the early diagnosis of haemostatic failure but also to guide replacement therapy during the fibrinopenic state.

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**Conflict of interest:** None declared

**Ethical approval:** The study was approved by the Institutional Ethics Committee

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