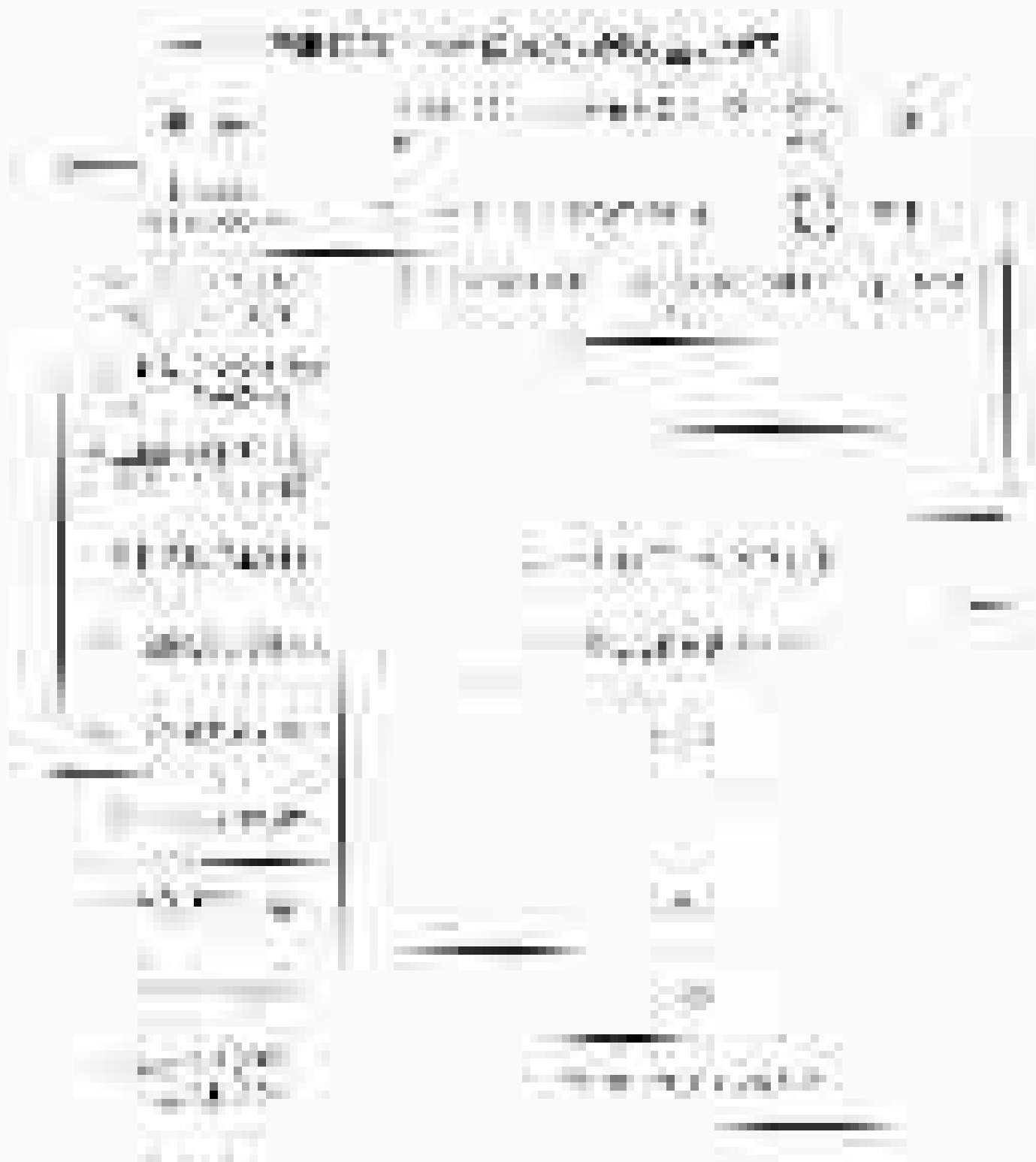


# 中国共产党第十七届中央委员会第五次全体会议 通过

## 中共中央关于制定国民经济和社会发展第十二个五年规划的建议



中国共产党  
中央委员会



## 四

此處有兩處，一處是說「我」，另一處是說「我」。這兩處的「我」，都是指「我」自己，不是指「我」的「我」。這兩處的「我」，都是指「我」自己，不是指「我」的「我」。

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the first time. — I hope you will be  
able to get some good news from  
the U.S. about the new law. —  
I am sending you a copy of the  
newly published "Principles of  
Economics" by Prof. Frank G.  
Loyd, which is a very good book.  
It is published by the McGraw-Hill  
Book Company, New York, and  
is now in its second edition. —  
I am sending you also a copy of  
the "Principles of Economics" by  
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New York, and is now in its second  
edition. —

1. **What is the primary purpose of the study?**  
The primary purpose of the study is to evaluate the effectiveness of a new treatment for depression compared to a placebo. The study will also assess the safety and side effects of the new treatment.

2. **Who is eligible to participate in the study?**  
Eligible participants include adults aged 18-65 years old who have been diagnosed with major depressive disorder. Participants must be willing to take part in weekly visits and follow-up assessments.

3. **What are the inclusion criteria for the study?**  
Inclusion criteria include having a diagnosis of major depressive disorder, being 18-65 years old, and being willing to participate in the study.

4. **What are the exclusion criteria for the study?**  
Exclusion criteria include having a history of suicidal behavior, being pregnant or nursing, having a history of substance abuse, and having other medical conditions that would interfere with participation in the study.

5. **How long will the study last?**  
The study will last approximately 12 months, including a 4-month baseline period, 8 months of treatment, and 2 months of follow-up.

6. **What are the treatment options available in the study?**  
Participants will receive either the new treatment or a placebo. Both treatments will be administered orally once daily. The new treatment is a combination of two medications, while the placebo is a dummy pill.

7. **What are the potential risks and benefits of participating in the study?**  
Potential risks include side effects from the medication, such as nausea, drowsiness, and headache. Benefits include the possibility of receiving effective treatment for depression and contributing to medical research.

8. **What are the costs associated with participation in the study?**  
There are no costs associated with participation in the study, as it is funded by the pharmaceutical company.

9. **What happens if I decide to withdraw from the study?**  
If you decide to withdraw from the study, you will be free to do so at any time. You will still be able to receive treatment for your depression, and your information will be kept confidential.

10. **What happens if I experience an adverse event during the study?**  
If you experience an adverse event, you should immediately contact the study team. They will assess the situation and provide appropriate care.

