EFFECT OF PROPOLIS EXTRACT AND PROPOLIS CANDIES ON STREPTOCOCCUS SOBRINUS GROWTH

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INTRODUCTION
Conventionally, herbal medicines have been used in Indonesia because their ingredients are relatively safe compared to synthetic drugs [1]. Indonesia has abundant natural resources that are easy to obtain—more than 9609 species of plants that can be sourced for herbal medicines [2]. One ingredient found in Indonesia that may inhibit bacteria colonization is propolis [3]. Flavonoid is believed to be a component of propolis [4]. The antibacterial properties of flavonoids have been proven by the previous studies; for example, the studies found that propolis might decrease Staphylococcus aureus and Streptococcus mutans bacteria inside the oral cavity [4,5]. Honey is another herbal ingredient with antibacterial properties. Its wound healing, infection-fighting, and inflammation-reducing abilities have been tested experimentally and clinically in the medical field. Like propolis, honey contains a bioactive ingredient (similar to flavonoid) that is responsible for its antibacterial effect.

In general, oral disease can be divided into two categories: Soft and hard tissue. The most common hard tissue disease is dental caries. The etiology of dental caries is highly influenced by plaque. Dental plaque is an oral biofilm that grows on the surface of teeth. Although the flora of the oral cavity is varied and complex, two species of streptococcus are most commonly associated with dental caries: S. mutans and Streptococcus sobrinus [6]. Propolis has been developed into an oral health product that comes in capsule, liquid, toothpaste, and mouthwash forms. In this study, propolis was packed into candies to make consumption more practical. Previous studies proved that honey propolis candies decrease the prevalence of S. mutans in the oral cavity. Therefore, the inhibitory effect of propolis extract and propolis candies on S. sobrinus growth was tested in this study.

METHODS
Making the S. sobrinus bacteria solution
A colony of S. sobrinus bacteria was transferred from a Petri dish into a brain heart infusion (BHI) medium. The centrifuge tube containing the BHI and S. sobrinus culture was incubated for 1×24 hrs in an anaerobic environment.

Making the propolis extract solutions
Concentrations of 0.5%, 1%, 5%, 10%, and 20% propolis solution were used in this study. A 20% glycerin solution was used to dilute the propolis extract to obtain the desirable concentrations. The propolis extracts were filtrated using a 0.22 µL syringe filter. The extract solutions were stored inside a 4°C refrigerator until their use.

Making the candy solution
First, propolis honey sucrose candies, propolis honey palm sugar candies, and X propolis candies were weighed. Then, the candies were mashed into small pieces using a mortar and pestle. The three types of candies were crushed up individually, into 3 separate mixtures. The candy powder was then transferred into an Erkenmeyer that contained 10 mL of BHI solution. Next, the candy solution was filtrated using a 0.22 µL syringe filter and stored inside a 4°C refrigerator until its use.

Determining the minimum inhibitory concentration (MIC) and minimum bactericidal concentration (MBC)
The S. sobrinus bacteria suspension was transferred into 96-well plates in triplo; either 100 µL of propolis extract solution or no propolis extract (the control) was then added into the 96-well plate. The samples were incubated for 1×24 hrs in an anaerobic environment. Afterward, the optical density reading was taken using an ELISA reader with a 450 nm wave length. The formula for the MIC is shown below:

\[ \text{Inhibition} \% = 1 - \left( \frac{\text{OD sample} - \text{OD blank}}{\text{OD control} - \text{OD BHI}} \right) \]

The MIC was determined based on the inhibition number reaching ≥ 90%. The sample result was placed in 10 µL of BHI agar. The lowest concentration of propolis extract with zero bacterial growth on the BHI agar was designated the MBC.

Quantitative bacteria test after candy exposure
The S. sobrinus suspension was transferred into a 96-well plate in 100 µL increments. Each bacteria-containing well was exposed to one of the four: 100 µL of the propolis honey sucrose sweetener candy
solution, propolis honey palm sugar candy solution, X propolis candy solution, and control (without propolis solution). The samples were incubated for 1×24 hrs in an anaerobic environment. Next, 10 µL of each microplate sample was transferred by Eppendorf pipette to solid agar and smeared using a wire loop. The samples were then incubated for 1×24 hrs at 37°C. The colony-forming units (CFU) on the surface of the BHI agar were calculated using the standard plate count (SPC) method.

RESULTS

The results of the propolis extract’s inhibition of *S. sobrinus* are shown in Table 1.

The inhibition mean value of the 0.5% propolis extract was 58% (±2.51) and the *p*-value, obtained through comparison to the control group, was <0.025. MIC is determined from inhibition values of ≥90%. As shown in Table 1, the first propolis extract solution to reach 90% was the 5% concentration. Table 2 shows calculations of the *S. sobrinus* colonies exposed to propolis extract.

As shown in Table 2, the 0.5% propolis extract *S. sobrinus* colony concentration was 48×10^5 CFU/mL. In the 1% and 5% propolis extracts, the *S. sobrinus* colony concentrations were 23×10^5 CFU/mL and 9×10^5 CFU/mL, respectively. No bacteria colonies grew in the BHI agar for the 0.5%, 15%, or 20% propolis extract concentrations. The lowest propolis extract concentration that had no *S. sobrinus* colony, the 10% concentration, was designated the MBC. Therefore, the MBC score of the propolis extract was 10%. Table 3 shows the bacterial colony counts after exposure to honey propolis palm sugar candies, honey propolis sucrose sweetener candies, and X propolis sugar candies.

As shown in Table 3, the lowest bacterial colony count occurred after exposure to propolis palm sugar candies (151×10^2 CFU/mL). The ANOVA showed a significant difference (*p*<0.025) between the propolis palm sugar candy group and the control group. *S. sobrinus* colonies also decreased in the honey propolis sucrose sweetener candy (208×10^2 CFU/mL) and X propolis candy (345.5×10^2 CFU/mL) groups. However, a one-way ANOVA test between these groups showed no significant difference (*p*<0.025).

### Table 1: Inhibition values of propolis extracts against *Streptococcus sobrinus* bacteria

| Propolis extract and control group concentration (%) | Inhibition value (%) | Inhibition mean value (%) ± SD | p-value against control group (no propolis extract) |
|-----------------------------------------------------|----------------------|-------------------------------|---------------------------------|
| Control group (no propolis extract)                 |                      |                               |                                 |
| 0.5                                                 | 0                    | 0                             | 0.000*                          |
| 1                                                   | 58                   | 56                            | 61                              |
| 5                                                   | 71                   | 71                            | 72                              |
| 10                                                  | 93.5                 | 93.8                          | 94.3                            |
| 15                                                  | 93.9                 | 94.2                          | 94.4                            |
| 20                                                  | 47                   | 42                            | 43                              |

There were significant differences (*p*<0.025), SD: Standard deviation

### Table 2: *S. sobrinus* colony calculations after propolis extract exposure

| Propolis extract and control group concentration | Number of *Streptococcus sobrinus* colonies (10^5 CFU/mL) |
|--------------------------------------------------|-----------------------------------------------------------|
| Control group (no propolis extract)              | 780                                                       |
| 0.5                                              | 48                                                        |
| 1                                                | 23                                                        |
| 5                                                | 9                                                         |
| 10                                               | 0                                                         |
| 15                                               | 0                                                         |
| 20                                               | 0                                                         |

*S. sobrinus*: *Streptococcus sobrinus*

DISCUSSION

Based on the results, it appears that the flavonoid properties in propolis inhibit the growth of *S. sobrinus*. Flavonoid destroys the cytoplasmic membrane, causing leakage of the cell contents [1]. Propolis most effectively inhibits the growth of *S. sobrinus* during cell division. During this process, the thin surface layer allows the flavonoid to easily penetrate the cell wall and damage the cell’s contents [7]. Propolis’ phenol properties may also inhibit the growth of *S. sobrinus*. These compounds bond to bacterial proteins. A weak bond between the protein-phenol forms in low concentrations, allowing phenol penetration into the cells, which denatures the protein and inhibits bacterial growth. In high concentrations, phenol can cause protein coagulation and lysis of the cell membrane. Barrientos et al. (2013) stated that the active substance in propolis depends on the compound’s species and collection location [8]. Propolis extracts with higher levels of active substances can more effectively inhibit bacteria growth [8]. The local propolis used in this study had 0.26% flavonoid, whereas Turkish propolis has a flavonoid content of 1.35%.

This study’s test results showed that the 0.5-10% propolis concentrations had increasing values of inhibition. However, the propolis extract’s inhibition against the *S. sobrinus* declined at 5% and 20% concentrations. These results differed from the BHI agar colony counts, where the higher concentrations of propolis extract resulted in the growth of fewer colonies. The inhibition value in this study was obtained from the optical density sample. Several factors determine the reading of optical density, such as consistency and specimen color [8]. In this study, the 15% and 20% concentration solutions of propolis extract had more consistency and concentrated color than the 0.5%, 1%, and 10% concentrations. This led to increased optical density for the 15% and 20% concentrations. Optical density values reflect the accumulation of live and dead bacteria [9]. Hence, it is possible that the solutions with the highest optical density had no existing live bacteria. Selkoe et al. stated that when assessing the growth of bacteria, the optical density reading is not always identical to the calculation results of bacteria on agar [10]. Therefore, a SPC method was used in this study to control the optical density value.

This study results showed that honey propolis palm sugar candies, honey propolis sucrose sweetener candies, and X propolis candies all decreased the colonization of *S. sobrinus* bacteria compared to the control group. The bacterial colonies for the honey propolis palm sugar candy test group decreased more significantly than the colonies for the honey propolis sucrose sweetener candy and X propolis candy groups. It can be concluded that honey propolis palm sugar candy and propolis honey sucrose sweetener candy. Carbohydrate intake contributes to the growth of *S. sobrinus*. Previous study stated that streptococci bacteria grow quickly because they can utilize various types of carbohydrates as additional nutrients for metabolism and cell growth [11]. This occurs because *S. sobrinus* can produce enzymes that catalyze hydrolysis. Sucrose is a carbohydrate utilized by *S. sobrinus* in the process of fermentation. Fewer *S. sobrinus* colonies were present
The propolis candies contained 5% propolis extract $9 \times 10^5$ CFU/mL of colonized $S. \ sobrinus$ was observed after exposure to the 5% propolis extract, whereas the mean $S. \ sobrinus$ population after consecutive exposures to honey propolis palm sugar and honey propolis sucrose sweetener candies was $131 \times 10^5$ CFU/mL and $208 \times 10^5$ CFU/mL, respectively. This result shows that propolis extract more effectively prohibits the growth of $S. \ sobrinus$ bacteria than honey propolis candy. The sugar content of the honey propolis candy likely lessened its prohibitive effect.

**CONCLUSIONS**

This study concludes that propolis extract may inhibit the growth of $S. \ sobrinus$ with a 5% MIC value and 10% MBC. Propolis honey sugar palm candy more effectively prohibits $S. \ sobrinus$ bacteria colonization than propolis honey sucrose sweetener candy and X propolis candy. More research is needed regarding the components contained in the propolis honey candy, particularly palm sugar and sugar and how each affects bacterial growth.

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Table 3: Mean bacterial colony count after exposed with propolis candies

| Propolis candy and control group | Sample I | Sample II |
|-------------------------------|----------|----------|
| Control group (no propolis)   | 780      | 1550     |
| PA                            | 113      | 149      |
| PM                            | 201      | 215      |
| PX                            | 328      | 363      |
| Mean $Streptococcus\ sobrinus$ colony count±SD (10^5 CFU/mL) | 1115±403 | 131±25.45 |
| p-value against control group (without propolis) | 0.019* | 0.027 |

*There were significant differences (p<0.025). PA: Honey propolis palm sugar candies, PM: Honey propolis sucrose sweetener candies, PX: X propolis candies, $S. \ sobrinus$: Streptococcus sobrinus