

## LAMPIRAN

### A. Contoh responden

#### 1. Diagnosis awal (*pre-test*)



2. Hari ke-7



3. Hari ke-14 (*Pre-test*)

## B. Data Frekuensi dan Persentase Subjek pada Kelompok Kontrol

**Frequencies****Statistics**

|   |         | Pretestair | Posttestair | Posttestair2 |
|---|---------|------------|-------------|--------------|
| N | Valid   | 8          | 8           | 8            |
|   | Missing | 0          | 0           | 0            |

**Frequency Table****Pretestair**

|       |        | Frequency | Percent | Valid Percent | Cumulative Percent |
|-------|--------|-----------|---------|---------------|--------------------|
| Valid | ringan | 5         | 62.5    | 62.5          | 62.5               |
|       | sedang | 3         | 37.5    | 37.5          | 100.0              |
| Total |        | 8         | 100.0   | 100.0         |                    |

**Posttestair**

|       |   | Frequency | Percent | Valid Percent | Cumulative Percent |
|-------|---|-----------|---------|---------------|--------------------|
| Valid | 1 | 5         | 62.5    | 62.5          | 62.5               |
|       | 2 | 3         | 37.5    | 37.5          | 100.0              |
| Total |   | 8         | 100.0   | 100.0         |                    |

**Posttestair2**

|       |   | Frequency | Percent | Valid Percent | Cumulative Percent |
|-------|---|-----------|---------|---------------|--------------------|
| Valid | 1 | 5         | 62.5    | 62.5          | 62.5               |
|       | 2 | 3         | 37.5    | 37.5          | 100.0              |
| Total |   | 8         | 100.0   | 100.0         |                    |

## C. Data Frekuensi dan Persentase Subjek pada Kelompok Eksperimen

**Statistics**

|   |         | Pretestzam | Posttestzam | Posttestzam<br>2 |
|---|---------|------------|-------------|------------------|
| N | Valid   | 8          | 8           | 8                |
|   | Missing | 0          | 0           | 0                |

**Frequency Table**

|       |        | Pretestzam |         |               |                       |
|-------|--------|------------|---------|---------------|-----------------------|
|       |        | Frequency  | Percent | Valid Percent | Cumulative<br>Percent |
| Valid | ringan | 4          | 50.0    | 50.0          | 50.0                  |
|       | sedang | 1          | 12.5    | 12.5          | 62.5                  |
|       | berat  | 3          | 37.5    | 37.5          | 100.0                 |
|       | Total  | 8          | 100.0   | 100.0         |                       |

|       |        | Posttestzam |         |               |                       |
|-------|--------|-------------|---------|---------------|-----------------------|
|       |        | Frequency   | Percent | Valid Percent | Cumulative<br>Percent |
| Valid | ringan | 4           | 50.0    | 50.0          | 50.0                  |
|       | sedang | 1           | 12.5    | 12.5          | 62.5                  |
|       | berat  | 3           | 37.5    | 37.5          | 100.0                 |
|       | Total  | 8           | 100.0   | 100.0         |                       |

|       |        | Posttestzam2 |         |               |                       |
|-------|--------|--------------|---------|---------------|-----------------------|
|       |        | Frequency    | Percent | Valid Percent | Cumulative<br>Percent |
| Valid | ringan | 4            | 50.0    | 50.0          | 50.0                  |
|       | sedang | 1            | 12.5    | 12.5          | 62.5                  |
|       | berat  | 3            | 37.5    | 37.5          | 100.0                 |
|       | Total  | 8            | 100.0   | 100.0         |                       |

D. Data SPSS Penelitian Uji *Chi Square*

## Case Processing Summary

|  | Valid                 |         | Cases Missing |         | Total |         |
|--|-----------------------|---------|---------------|---------|-------|---------|
|  | N                     | Percent | N             | Percent | N     | Percent |
|  | airzamzam * turunlesi | 16      | 100.0%        | 0       | 0.0%  | 16      |

## airzamzam \* turunlesi Crosstabulation

|           |                    | turunlesi          |        | Total  |       |
|-----------|--------------------|--------------------|--------|--------|-------|
|           |                    | 1                  | 2      |        |       |
| airzamzam | 1                  | Count              | 8      | 0      | 8     |
|           |                    | % within turunlesi | 66.7%  | 0.0%   | 50.0% |
| 2         | Count              | 4                  | 4      | 8      |       |
|           | % within turunlesi | 33.3%              | 100.0% | 50.0%  |       |
| Total     | Count              | 12                 | 4      | 16     |       |
|           | % within turunlesi | 100.0%             | 100.0% | 100.0% |       |

Chi-Square Tests<sup>d</sup>

|                                    | Value              | df | Asymptotic Significance (2-sided) | Exact Sig. (2-sided) | Exact Sig. (1-sided) | Point Probability |
|------------------------------------|--------------------|----|-----------------------------------|----------------------|----------------------|-------------------|
| Pearson Chi-Square                 | 5.333 <sup>a</sup> | 1  | .021                              | .077                 | .038                 |                   |
| Continuity Correction <sup>b</sup> | 3.000              | 1  | .083                              |                      |                      |                   |
| Likelihood Ratio                   | 6.904              | 1  | .009                              | .077                 | .038                 |                   |
| Fisher's Exact Test                |                    |    |                                   | .077                 | .038                 |                   |
| McNemar Test                       |                    |    |                                   | .125 <sup>c</sup>    | .063 <sup>c</sup>    | .063 <sup>c</sup> |
| N of Valid Cases                   | 16                 |    |                                   |                      |                      |                   |


a. 2 cells (50.0%) have expected count less than 5. The minimum expected count is 2.00.

b. Computed only for a 2x2 table


c. Binomial distribution used.

d. For 2x2 crosstabulation, exact results are provided instead of Monte Carlo results.

## E. Form Etik



**FAKULTAS KEDOKTERAN UNIVERSITAS ISLAM BANDUNG**  
**KOMITE ETIK PENELITIAN KESEHATAN**  
 Jl. Tamansari No. 22 PO.BOX 1357 Telp. (022) 4203368 (hunting) Pes. 6905 Fax. 4231213 Bandung 40116



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**PERSETUJUAN ETIK**  
**ETHICAL APPROVAL**

Nomor: 52/Komite Etik.FK/IV/2019

*Bismillahirrahmanirrahim*

Komite Etik Penelitian Kesehatan Fakultas Kedokteran Universitas Islam Bandung, dalam upaya melindungi hak asasi dan kesejahteraan subjek penelitian kesehatan serta menjamin bahwa penelitian yang menggunakan formulir survei/registrasi/surveilans/epidemiologi/humaniora/sosial budaya/ bahan biologi tersimpan/sel punca dan nonklinis lainnya berjalan dengan memperhatikan implikasi etik, hukum, sosial, dan nonklinis lainnya yang berlaku telah mengkaji dengan teliti proposal penelitian berjudul:

*The Health Research Ethics Committee, Faculty of Medicine, Universitas Islam Bandung in order to protect the rights and welfare of the health research subject, and to guaranty that the research using survey quetionnaire surveillance epidemiology/humanities social-cultural archived biological materials/ stem cell other non-clinical materials, will carried out according to ethical, legal, social implications and other applicable regulations, has been troughly reviewed the proposal entitled:*

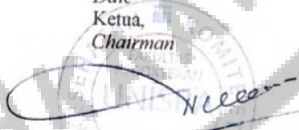
**Efek Pemberian Air Zamzam terhadap Perbaikan Derajat Akne Vulgaris pada Mahasiswa Fakultas Kedokteran Unisba**

|                                      |   |
|--------------------------------------|---|
| Nama mahasiswa<br><i>Student</i>     | : Gaby Syafira Siti Halimatussadiyah            |
| NPM<br><i>Student Batch Number</i>   | : 10100116050                                   |
| Pembimbing 1<br><i>Supervisor 1</i>  | : Anita Indriyanti,dr., M.Kes                   |
| Pembimbing 2<br><i>Supervisor 2</i>  | : Nurul Romadhona, dr., MMRS                    |
| Nama institusi<br><i>Institution</i> | : Fakultas Kedokteran Universitas Islam Bandung |

penelitian tersebut dapat disetujui pelaksanaannya.  
*hereby declare that the proposal is approved.*

Demikian, surat keterangan ini dibuat dengan sebenar-benarnya dan untuk digunakan sebagaimana mestinya.

Ditetapkan di: Bandung  
*Issued in*  
 Pada tanggal: 30 April 2019  
*Date*  
 Ketua,  
*Chairman*



**Prof. Herry Garna, dr., Sp.A(K), Ph.D.**

Keterangan/notes:  
 Persetujuan etik ini berlaku selama satu tahun sejak tanggal ditetapkan.  
*This ethical clearance is effective for one year from the due date.*  
 Pada akhir penelitian, laporan pelaksanaan penelitian harus diserahkan ke Komisi Etik Penelitian Kesehatan.  
*In the end of the research, progress and final summary report should be submitted to the Health Research Ethics Committee.*  
 Jika ada perubahan atau penyimpangan protokol dan/atau perpanjangan penelitian harus mengajukan kembali permohonan kajian etik penelitian.  
*If there be any protocol modification or deviation and/or extension of the study, the principal investigator is required to resubmit the protocol for approval.*  
 Jika ada kejadian serius yang tidak diinginkan (KTD) harus segera dilaporkan ke Komisi Etik Penelitian Kesehatan.  
*If there are serious adverse events (SAE) should be immediately reported to the Health Research Ethics Committee.*

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