Development and Cultivation of Local Kidney Bean 
(*Phaseolus Vulgaris* L) Through Breeding to use Multi gamma Irradiation Technique (Nuclear)

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**Abstract:** Mutation on the all species of plant were generally caused by gamma or multi gamma radiation sources. The largest effect of that was occurred on chromosome and genetic factor, specially on structure and composition of chromosome and Deoxyribo Nucleic Acid (DNA). This case can be used for breeding of several important plants in Agricultural science. The objectives of this research is to develop of local kidney bean (*Phaseolus Vulgaris* L) that tolerant to dry condition and high production. Multi gamma radiation and selection are the main methods to use in this research and other methods comprised of observation/survey, sampling, comparison, analyzing and interpretation. The number of mutant varieties of kidney bean (superior seed of mutant) obtained in this research based on selection method are ten mutant varieties. The growth percentage rate, protein content of control and mutant variety of kidney bean were estimated respectively of 97, 22.54 and 24.21%. The production of mutant kidney bean was between 3.97 tons/ha up to 5.28 tons/ha and the average production was 4.71 tons/ha. The average production of control was 2.60 tons/ha, in order that the production was significantly increased by 44.80%.

**Keywords:** Breeding, Kidney Bean, Multigamma, Irradiation

**Introduction**

Mutation is a spontaneously changing process which occurred on genetic matter of an organism (called spontaneous mutation) and random and also through induce mutation (Soedjono, 2003). The abnormality on plant and other organism was irradiated by gamma or multi sources show the changing on genetic level or deoxyribo nucleic acid (DNA) and chromosome. Those process produce variations of new genetic as the basic of plant selection (natural or breeding), in order that, the breeder is easy to select of genotive apropriate to purpose of breeding (Gepts and Hancock, 2006; Agusrial, 2008; Carsono, 2008). On general, physical mutagent is high energy which produced by nuclear reaction from radioactive sources.

Theory of radiation effect there are two categories (Hollaender, 2002) i.e.: (1) Target theory or direct action theory. The biologists qualitatively and quantitatively investigate and explain the cell multiplication and mutation on the organism and around vital structure or molecular structure. They begin their investigation on outher morphology effect of organism, continuously to sensitive spot and vital structure (Handayani, 2009). The continuously research on citology and genetics obtained the changing in molecular structure. (2) Indirect action theory. This theory comes from chemicalists whose begin their investigation about radiation effect on act aspect of molecular and continued to macromolecular which cell composing, like as DNA, RNA, protein, etc for easy explaining of organism mutation.

**Radiation Effect Ionizes Nucleic-Acid and Nucleo-Protein**

The important components in chromosome support characterisric of generation are Nucleic-acid and nucleo-protein (Hollaender, 2002). The aberation of chromosome is caused by ionization energy disturbs of cell fission and mutaton. Multi gamma radiation produces depolarization and viscosity descent on Thymonucleic Acid (TNA), impedes syntesis of Deoxyribo Nucleic Acid (DNA). The several approaches...
of physics and biology have been done for illustrating of dosage limit and speed of low dosage. According to microdosimetry aspect, low dosage is smaller than 1 mGy, radiobiology: Low dosage is 20 mGy, epidemiology: Low dosage 200 mGy (UNSCEAR, 2005).

**Radiation effect on DNA and Chromosome**

DNA structure formed of double helicks which composed from bundle between phosphate group and dioxiribo sugar that form of strand DNA and bundle between nitrogen bases, which connect to two strands DNA. A large parts of DNA broken are occurred on bases broken, bases lost, the bundle between bases has broken and the bundle of sugar and phosphate has broken, bases lost, the bundle between bases has broken and the bundle of sugar and phosphate has broken, in order that, occurred broken on one strand is called single strand break (ssb). This damage can be quickly reconstructed without mistake by enzymatic repairs process with using strand DNA that is not break as mold. Cell can do the contraction process to the DNA broken in a few hours, but can be not perfect, mainly to the broken of DNA is called double strands breaks (dsb) (Brenner et al., 2006). The reconstruction process with mistaking causes mutaion of abnormality genetics and chromosome. The changing of chromosome structure is caused by gamma or multigamma radiation. Normally, chromosome comprises of upper-arm and fore-arm connected by a centromer. Multigamma radiation causes forming of: (1) assentric fragment (formed of chromosome fragment without centromer), (2) disentric chromosome (chromosome has two centromers), (3) ring chromosome, (4) translocation (removal of genetic matter betwen chromosome arm) (UNSCEAR, 2005).

According to IAEA statement (IAEA, 2008), mutation on the plant is spontaneously changes of genetic matters in cells caused by: (1) rearrangement occurred on chromosome structure, (2) changing in genetics, (3) segments duplication of chromosome loss. Radiation technique has several superiority among others: (1) radiation technique is easy to do and practical, (2) the change of genotive a few only, but causes much changes of characteristics on generation species, (3) the generation species obtain in the short time.

Dose standard of gamma/multi gamma radiation is used on breeding of plants (IAEA, 2003; 2004; 2006): (1). Mutation on plants: 100 rads up to 3,000 rads, (2). Mutation in seeds plant: 1,000 rads up to 4,000 rads, (3). Growth stimulation of seeds plant: 250 rads up to 1,000 rads, (4). Growth obstruction on root: 5,000 up to 10,000 rads (NNEA, 2005). This research focused on development of local kidney bean from Manggarai Flores Indonesia with using multi gamma irradiation technique. In this method was obtained several variations of mutant, in order to easily selection of superior variety. The general characteristics of mutant of kidney bean variety are: (1) high production, (2) tolerant to dry conditions (Hartati, 2000), (3) the age of mutant is shorter, (4) tolerant to germ specially viruses (Radiyanto et al., 2011), (5) tolerant to plant disease like as Henose-pilachna signstipennis or Epilacha signatipennis, Etilla zickenella, Agromointa phaseoli, Aphis gossiphi, Phytonecta signata and P. calcites, Lamprosema indicata and L. diamenalis (Purana and Mitra, 2008), (7) the quality of seed increase (content of protein and fat) (Irwan, 2006; NNEA, 2008). The development of local kidney bean in these research, uses multigamma radiation techniques. Multigamma radiation techniques lead of genetic effect such as the changes of structure and composition of chromosome and molecule of Deoxiribo Nucleat Acid (DNA) on several species of food plant.

Kidney bean is one kind of legumes that can grow at tropic and sub-tropic lowland up to altitude 2,000-2,500 m from sea level on temperatures 16 up to 27°C and rain fall 900 mm up to 1,500 mm, air humidity 50 up to 60% (Rukmana, 2003). The average production of kidney bean until 2008 on national level was very low (only 1.2 tons/ha up to 2.13 tons/ha), just the opposite that the demand of consumer increase every year (Purana and Mitra, 2008). The number of pods per tree is 8 up to 10 pods, the number of seeds per pod is 1 up to 3 and high plant is 15 cm up to 20 cm. Kidney bean content of protein (23,15%), folic acid, vitamin-B, folasin, tiamin, calcium, phosphor, iron and complex carbohydrate. The content of complex carbohydrate and fiber in kidney bean can decrease concentration of cholesterol in the blood (Purana and Mitra, 2008; UNSCEAR, 2005).

The comodity production of kidney bean per hectarre in Indonesia is not achieve maximum. That is influenced by soil factor, which damaged and is poor of micro-elements, growing hormone, dry conditions, germ, climate and the using of superior seeds (Purana and Mitra, 2008). These research aimed: To develop local kidney bean variety from Manggarai Flores Indonesia through breeding with aplication of multigamma radiation technique and carefully selection to obtain superior seed with high production and tolerant to dry condition.

Since 2010, researchers succeed to develop erect local penaut and creep local peanut from East Sumba with using multi gamma radiation. The increase of mean production 43.86% for erect local peanut and 42.22% for erect local peanut, or mean production 5.7 tons/ha (from 3.2 tons/ha up to 5.7 tons/ha) for erect local peanut, mean production potential 5.9 tons/ha and 4.5 tons/ha (from 2.6-4.5 tons/ha) for erect local peanut, production potential 4.7 tons/ha (Pasangka and Jaelani, 2011). Since 2011, continuous research obtained production to be revolved between 4.75 tons/ha up to 6.84 tons/ha for creep local peanut and between 3.95 tons/ha up to 5.45 tons/ha for erect local peanut (Pasangka and Jaelani, 2011).
Since 2012 up to 2014, researchers succeed to develop local soybean from Bajawa Flores Indonesia. The results of research are superior seed of mutant soybean with high production and tolerant to dry condition. There were ten varieties of superior seed of mutant were obtained from mutation by multi gamma radiation treatment. The production of mutant soybean (treatment sample) was 3.78 tons/ha up to 4.92 tons/ha, with mean production of 4.41 tons/ha. The mean production of control sample (initial soybean) was 2.54 tons/ha, with a significant increase of 42.40% (Pasangka and Refly, 2013).

Materials and Methods

Material Studied

The main instruments (Lab nuclear Physics Faculty of Sciences and Technology Nusa Cendana University) used in this research consist of: (1) Irradiator (Multi gamma radiation source: Co-60, Cs-137, Cd-109, Mn-54 and Sn-113 in one kit), (2) counter of radiation dose (Radiolet-50 gamma counter type), (3) protein analyzer, (4) tractor, (5) other equipments. The sample for breeding is a local kidney bean from Manggarai Flores Island Indonesia with two groups: Control and treatment samples.

Description Area

The area of research is located in Kupang West Timor Island Indonesia, at five areas (the name of locations are: Fukdale, Naibonat, Baumata, Taibenu and Bakunase). All of those areas have the same conditions such as the high level of salt and calcium, dry condition, soil structure and illumination. The level of salt and calcium respectively are 2.86 and 22.62%. The position of the area between 10°07’ 03.6” south latitude, 123°48’48.1” east longitude and 10°07’ 03.6” south latitude, 123°49’ 57.4” east longitude on 96 m altitude (96 m from sea level). The average illumination of the area is 5 h per day.

Methods

The main methods of research consist of: Observation/surveying, sampling, radiation, selection, comparison and interpretation. Collecting and data analysis are done with observation, measurement, protein analysis on initial kidney bean and also on treatment sample (mutant). Quality control is done to compare between physical and chemical characteristics of control sample (initial kidney bean) and treatment sample (mutant) that obtained from observation, measurement and calculation.

Procedures of Research

The procedures of this research consist of: (1) to choose research location and sample of local kidney bean, (2) to prepare research location, (3) to irradiate samples of kidney bean seeds as long as one hour, (4) to plant sample of kidney bean in the prepared area. The conditions of prepared area are specified to be based on data from department of agricultural, i.e.: The level of salt and calcium respectively are 2.86 and 22.62%, (5) watering if it is necessary, (6) weeding and cultivating, (7) observation to tenacity of germ, growth in dry area, high calcium and salt and physical characteristics which were needed for a standard comparison i.e., growth time, flowered age, tenacity of germ, adaptation, growth percentage, mass per 1,000 seeds, maximum production potential, mean production and also select plants. On the resemble harvest, is done selection, measurement of high plant, in time after harvest is measured of mass for a group of 1,000 kidney bean seeds, (8) to analyze protein (service analysis model), (9) drying and selection. The drying conditions is take time from 06.30 a.m until to 9,30 a.m, (10) For quality control is done comparison between physical and chemical characteristics of control sample (initial kidney bean) and treatment sample (mutant), that obtained from observation/survey, measurement and calculation, (11) The last procedure is to put insecticide sufficient to mutant (superior seeds of kidney bean), so to be spared from pest and or germ and storage for continuously development. The first selection of kidney bean plant is done since the age of plant is one month, the second selection since the age of plant two months, M_{0} selection since near to harvest and the third selection after harverst.

Observe and Measures

The amount of physical characteristics of kidney bean (Control sample and mutant) during growth and after harvest was observed and measured like as adaptation, tenacity of germ, grow time, flower age, the length of plant and weight per 1,000 seeds, protein content, potential production and mean production. In this case were selected 100 plants of each variety for measuring. Table 1 contain any informations about the important physical and chemical characteristics of kidney bean and Table 2 special contain of Data for evaluating of growth percentage rate (the number of seeds was not grown on the control sample and treatment sample). The calculation result of data in Table 3 also included in Table 1.

Research Design

The samples of local kidney bean were chosen with two groups, i.e., control (initial kidney bean) and treatment samples. The treatment sample (seeds of kidney bean) was irradiated by multigamma sources on radiation dose 3.500 rads. The samples were planted in...
the same time and location. The time planting is started from 06.00 a.m until to 09.00 a.m and all locations have the same conditions like as level of salt and calcium, dry condition, the same altitude (96 m from sea level). Three plant selections were done according to the research design on Fig. 1. The first selection of kidney bean plant was done since the age of plant was one month, the second selection since the age of plant was two months, M0 selection since near to harvest and the third selection after harvest.

![Research design](image)

Fig. 1. Research design

| Description                  | Control sample of kidney bean (Initial Variety) | Mutant kidney bean (Treatment sample or generation Variety) |
|------------------------------|--------------------------------------------------|-----------------------------------------------------------|
| Grow time                    | 9 days after planted (dap)                       | 5 dap                                                     |
| Flowered age                 | 68 dap                                           | 45 dap                                                    |
| The average high of plant    | 41.23 cm                                         | 67.40 cm                                                  |
| Tenacity of germ             | Not tenacity                                     | Tenacity                                                  |
| Adaptation                   | Adaptation to area with high calcium and salt.   | Adaptation to area with high calcium and salt, dry condition. |
| Growth percentage rate       | 80%                                              | 97%                                                       |
| The age of plant             | 105 days after planted (dap)                     | 73 days after planted (dap)                               |
| Protein content              | 22.54%                                           | 24.21%                                                    |
| The number of pods per tree  | 4-10 (average: 6)                                | 5-22 (average: 16)                                        |
| The number of seeds per pods | 1-4                                              | 2-7                                                       |
| Mass per 1,000 seeds         | 374.88 grams                                     | 543.70 grams                                              |
| Production range             | 2.46 tons/ha - 2.81 tons/ha                      | 3.97 tons/ha - 5.28 tons/ha                               |
| Maximum production potential | 2.81 tons/ha                                     | 5.28 tons/ha                                              |
| The average production       | 2.60 tons/ha                                     | 4.71 tons/ha                                              |
Table 2. Production level at several planting locations of kidney bean (Control and treatment samples)

| Planting location (1 hectare) | Control sample | Treatment sample (Mutant) |
|-------------------------------|----------------|--------------------------|
|                               | Leaf ranges (stems) | Production level (tons/ha) | The average production (tons/ha) | Leaf ranges (stems) | Production level (tons/ha) | The average production (tons/ha) |
| Fukdale (P1)                  | Adventiti-root      | 2.81                      | Adventiti-root                   | 5.28               |
| Naibonat (P2)                 | Ous root,           | 2.54                      | Ous root,                        | 4.96               |
| Baumata (P3)                  | 5-7 Large           | 2.52                      | 6-10 Large                       | 4.23               |
| Taibenu (P4)                  | Straight            | 2.69                      | Straight                         | 5.12               |
| Bakunase (P5)                 | Root               | 2.46                      | Root                             | 3.97               |

Table 3. Data for evaluating of growth percentage rate (the number of seeds was not grown on the control sample and treatment sample or mutant of kidney bean)

| Sample Group | Control Sample | Mutant (treatment sample) |
|--------------|----------------|----------------------------|
|              | Total number of seeds every group | Number of seeds was not grown | Number of seeds was not grown |
| I            | 100            | 18                         | 4                           |
| II           | 100            | 17                         | 1                           |
| III          | 100            | 24                         | 5                           |
| IV           | 100            | 19                         | 2                           |
| V            | 100            | 22                         | 1                           |
| Mean         | 100            | 20                         | 3                           |

Statistical and Data Analysis

Statistical formula was needed to calculate of growth percentage rate and percentage of increasing of mean production. For testing of growth percentage rate, we choose five samples groups at random on control and treatment sample. The number of test sample is 100 seeds on every group. The number of seeds sample was not grown to be observed. The growth percentage rate was calculated by equation (Pasangka, 2015):

\[
GP = \left(\frac{T_{SG} - N_{SN}}{T_{SG}}\right) \times 100\% \tag{1}
\]

Where:

- \(GP\) = Growth percentage rate (%)
- \(T_{SG}\) = Total number of seeds to be planted (prepared sample)
- \(N_{SN}\) = The number of seeds was not grown

Mean production was calculated by equation formula:

\[
M = \frac{P_1 + P_2 + P_3 + P_4 + P_5}{5} \tag{3}
\]

where, \(P_1, P_2, P_3, P_4, P_5\) are production at 5 locations.

Protein content is calculated by equation:

\[
PPC = \frac{(B - E) \times N \times NaOH \times 0.014}{m} \times 100 \tag{4}
\]

where:

- \(PPC\) = Percentage of protein content
- \(B\) = Blank
- \(E\) = Example
- \(m\) = Mass of sample of kidney bean (control sample and treatment sample).

Flowered rate percentage can be calculated by formula:

\[
FRP = \left(\frac{FACS - FATS}{FACS}\right) \times 100\% \tag{5}
\]

where:

- \(FRP\) = Flowered rate percentage
- \(FATS\) = Flowered age of treatment sample
- \(FACS\) = Flowered age of control sample
The increasing percentage of mass per 1,000 seeds can be calculated by equation:

$$P_n = \left( \frac{MPTS - MPCS}{MPTS} \right) \times 100\% \quad (6)$$

Where:
- $P_n$ = Increasing percentage of mass per 1,000 seeds
- $MPTS$ = Mass per 1,000 seeds of treatment sample
- $MPCS$ = Mass per 1,000 seeds of control sample

The increasing percentage of mean pods per tree is calculated by equation:

$$IPPP = \left( \frac{MPPTS - MPPCS}{MPPTS} \right) \times 100\% \quad (7)$$

Where:
- $IPPP$ = The increasing percentage of mean pods per tree
- $MPPTS$ = The number of pods on treatment sample
- $MPPCS$ = The number of pods on control sample

Calculating

Based on equation (1), growth percentage rate of seeds on control sample and mutan can be determined. Growth percentage rate of control sample:

$$GP = \left( \frac{T_{SG} - N_{SG}}{T_{SG}} \right) \times 100\% = \left( \frac{100 - 20}{100} \right) \times 100\% = 80\%$$

Growth percentage rate of mutant:

$$GP = \left( \frac{T_{SG} - N_{SG}}{T_{SG}} \right) \times 100\% = \left( \frac{100 - 3}{100} \right) \times 100\% = 97\%$$

The percentage of increasing of mean production was determined by Equation 2, i.e.:

$$P = \left( \frac{M_p - M_c}{M_p} \right) \times 100\% = \left( \frac{4.71 - 2.60}{4.71} \right) \times 100\% = 44.80\%$$

Mean production of control sample:

$$M_c = \frac{P_1 + P_2 + P_3 + P_4 + P_5}{5}$$
$$= 2.81 + 2.54 + 2.52 + 2.69 + 2.46$$
$$= \frac{12.1}{5} = 2.42 \text{ tons/ha}$$

Mean production of mutant (treatment sample):

$$M_p = \frac{P_1 + P_2 + P_3 + P_4 + P_5}{5}$$
$$= \frac{5.28 + 4.96 + 4.23 + 5.12 + 3.97}{5} = 4.71 \text{ tons/ha}$$

Protein content:

$$PPC = \frac{(B - E) \times m \times N \times NaOH}{x} \times 100\%$$

Note: Protein content is direct calculated in protein analyzer as many as 22.54% for control sample (initial kidney bean) and 24.21% for treatment sample (mutant).

Flowered rate percentage:

$$FRP = \left( \frac{68 - 45}{68} \right) \times 100\% = 33.83\%$$

The increasing percentage of mass per 1,000 seeds:

$$P_n = \left( \frac{MPTS - MPCS}{MPTS} \right) \times 100\%$$
$$= \frac{543.70 - 374.88}{543.70} \times 100\% = 31.03\%$$

The increasing percentage of mean pods per tree:

$$IPPP = \left( \frac{16 - 6}{16} \right) \times 100\% = 62.50\%$$

Result and Discussion

Observes and Measures

The important physical and chemical characteristics were observed and measured of kidney bean on control sample and treatment sample (Mutant) included in Table 1 and the production level at the several planting locations of local kidney bean on control and treatment samples incuded in Table 2.

Figure 2a, 2c and 2e shows three examples of the physical growth of control samples (initial local kidney bean from Manggarai Flores Indonesia) and the physical growth of mutant (variety of kidney bean was obtained from mutation by multigamma radiation or treatment sample) were shown on Fig. 2b, 2d and 2f.

The Fig. 2 shown that the growth (Fig. 2a control sample and Fig. 2b treatment sample) and fruits between control and treatment sample is significantly differenced with an important amounts of fruits has been observed for the treatment sample (Fig. 2d) compared to the control sample (Fig. 2c) and it’s clearly that the Fig. 2f (treatment sample) has many fruits than Fig. 2e (control sample).
Fig. 2. Figure 2a-e show three examples of the physical growth of control samples and Figure 2b-f show three examples of the physical growth of mutant (result from multi gamma radiation), (a) Control sample, (b) Treatment sample (irradiated by 3500 rads), (c) Control sample, (d) Treatment sample, (e) Control sample, (f) Treatment sample.
Fig. 3. The seeds of control sample (seeds of initial kidney bean)

Fig. 4. Three examples of mutant seed varieties were obtained from mutation by multigamma radiation, (a) Mutant-1 variety of kidney bean, (b) Mutant-2 variety of kidney bean, (c) Mutant-3 variety of kidney bean
Table 4. The physical and chemical characteristics of the ten mutant varieties were observed and measured

| Description                  | M01  | M02  | M03  | M04  | M05  | M06  | M07  | M08  | M09  | M10  |
|------------------------------|------|------|------|------|------|------|------|------|------|------|
| High of plant (cm)*          | 64.17| 64.45| 65.78| 66.12| 67.25| 67.86| 68.19| 68.84| 70.21| 71.13|
| The age of plant (days)      | 68   | 68   | 70   | 72   | 73   | 75   | 75   | 76   | 76   | 77   |
| The number of pods per tree* | 22   | 18   | 18   | 18   | 15   | 15   | 15   | 14   | 13   | 12   |
| The number of seeds per pod* | 3    | 4    | 4    | 4    | 5    | 5    | 5    | 6    | 7    | 7    |
| Color of seed*               | Wine | Brown| Brown| Pink | Dark | Bright| Pink | Pink | Wine | colored |
| Mass per 1,000 seeds*        | 546.10| 545.80| 545.76| 545.24| 543.62| 543.30| 543.18| 542.36| 541.14| 540.50|
| Adaptation to dry condition* | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| Adaptation to germ*          | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |

Figure 3 shows one example of seeds of control sample and Fig. 4a up to Fig. 4c, show three examples of mutant (mutant seeds of kidney bean was obtained by mutation by multigamma irradiation with ten varieties). Mutant-1 up to mutant-3 on Fig. 4, were obtained by a carefully selection. Selection is done appropriate to several important physical and chemical characteristics like as production level, age of plant, mass of a group 1,000 seeds, high of plant, flowered age and other characteristics.

The number of seeds was not grown on the control sample and mutant were included in Table 3 and the physical and chemical characteristics of the ten mutant varieties were observed, measured and calculated were included in Table 4. The varieties of mutant M01 to M010, were called the first generation or M1. The second, the third, etc generation or M2, M3, ...,Mn will be obtained in the next research, for this reason will be carefully selected on the second, third, etc planted on groups M01, M011, M012, ...,M01n, M02, M021, M022, ...,M02n etc. The second research will be examined homogeneous of mutant and the third research will be examined multi location, homogeneous of mutant and tolerant to dry condition and germ.

Data in Table 3 show the five groups of control and treatment samples for evaluating of growth percentage rate. The average number of seeds was not grown on control and treatment samples respectively were twenty seeds and three seeds. This case shows that the calculating result of growth percentage rate of control sample and treatment sample were 80 and 97%.

Data in Table 4 show the physical and chemical characteristics of the ten mutant varieties were observed and measured. The calculation result based on data in Table 4, give the average: High of plant, age of plant, number of pods per tree, number of seeds per pod and mass per 1,000 seeds respectively were 67.40 cm, 73 days, 16 pods, 5 seeds and 543.70 grams.

The Time Growth of Seeds, Growth Percentage Rate, Flowered Age and other Characteristics

The seeds of control sample (initial kidney bean) need 9 days to grow and less required time for the treatment sample (mutant), which was estimated of 5 days after planted (dap) and since the age is 8 days, the growth of sprout is 97% and the age of harvest on treatment sample (mutant) is 73 dap. This case shows that the treatment samples (mutant) grow faster than control sample (without irradiation). So, can be proposed that the treatment sample (mutant) has age is the short time compared to control sample (105 dap). The control sample is flowered on 68 days after planted (dap) and flowered age of treatment sample (mutant) is 45 days after planted (dap). Thus, clearly show that the treatment sample is flowered faster than control sample (is faster than 33.83%). The control sample was adapted to area with high calcium and salt and the mutant varieties or treatment sample of kidney bean was adapted to area with high calcium, salt and dry condition. The growth percentage rate of control sample (initial kidney bean) is 80% and the treatment sample (mutant) is 97%.

The Range Production, Average Production, Increase of Mean Production and Adaptation, Fertilizing

The mean or average mass of 1,000 seeds of control sample (initial kidney bean) is 374.88 grams and mutant sample is 543.70 grams. The increasing percentage of mass per 1,000 seeds is 31.03%. The increasing percentage of mean pods per tree is 62.50%. The range production of control sample (initial kidney bean) is 2.46 tons/ha up to 2.81 tons/ha and treatment sample (mutant) is 3.97 tons/ha up to 5.28 tons/ha. The maximum production potential of control sample is 2.81 tons/ha, with average production 2.60 tons/ha and mutant variety is 5.28 tons/ha with average production 4.71 tons/ha. Those data show that physical and chemical characteristics of mutant are superior than control sample (initial kidney bean). All of these characteristics were clearly different. The ten varieties of primer seed or superior seed were obtained from mutation by multigamma radiation and carefully selection. The improved traits stably inherited in the later generation of those varieties of soybean (10 varieties) will be tested on continuously research. NNEA gives report that the mean production of kidney bean only 4.10 tons/ha and can be adapted to area with high calcium but was not adapted to dry conditions (NNEA, 2008). In other researches (Pasangka and Jaelani, 2011; 2013) on corn, peanut and...
soybean breeding, they found that the generations of those plants have the traits stably inherited up to 7 generations and after that the production to be more decreased. Agricultural land in Kupang Timor Island Indonesia was dominated by soil with high calcium, high salt and dry condition and also germ. It’s necessary to develop plants that will be able to adapt in those conditions. Figure 4a up to 4c, show three examples of mutant seeds of kidney bean. Kidney bean seeds on: Figure 4a, tolerant to germ, dry condition and area is high calcium and salt, Fig. 4b tolerant to germ, dry condition and area is high calcium, Fig. 4c, tolerant to germ, dry condition and area with high calcium. The control sample is not tolerant to dry conditions, only adapt to high calcium and salt.

Contain of Nitrogen in soil on research location was not enough for good growing of plant (17%). According to International standard, this value was low (< 10%) was very low, 10-21% was low, 22-51% was medium, 52-75% was high, > 75% was very high, Rukmana, 2003) Because of that, fertilizing was done when the age of plant (kidney bean) about 24 days, (before flowered) to use NPK (Nitrogen-Phosphorous-Potassium) and Urea. Concentration of Nitrogen in Urea was 46%. The quantity of fertilizing used was 100 kilograms/ha NPK and 100 kilograms/ha Urea. It was enough for good growing of plant. It was appropriate with previous research (Rukmana, 2003).

Tenacity to Germ

Based on observations from first growth until to harvest time or during growth of plant was shown clearly that growth of treatment sample of kidney bean (mutant) tenacity to germ. In this case is shown by soft leaf since growth of kidney bean. This argument clearly shown on Fig. 2c (control sample) and Fig. 2f (treatment sample or mutant). The plant on Fig. 2e has not fruit and it’s leaf wrinkled because attacked by germ. According to the research result from Radiyanto et al. (2011), the kidney bean is not tenacity to several germs like as Aphis glycine, Riptortus Linearis, Sclerotium Rolfsii, Phachyrizi Phakospora and Xanthomonas Phaseoli.

The Containing of Protein

Protein content of control sample and treatment sample (mutant) respectively were 22.54 and 24.21%. According to literature (previous research), protein content of control sample is 23.15% (Puryana and Mitra, 2008; UNSCEAR, 2005).

Root, Leaf, Soil Condition, Temperature, Rainfall and Humidity

Generally, kidney bean has the adventitious root and large straight root with shallow lateral branching. The root strengthen on the growth of plants, transport of water and the other elements to the all parts of plants.

The first joints of plant (nodus) on pre growth formed pair of single leaf (Fig. 2b) and continuous on the all joints formed compound of leaf with three leaves. The form of leaf was oval, thin and green color. Leaf surface has soft feather (trachoma) on the both side. The flowers growth on the armpit of compound leaf stalks.

Based on observation result and previous research, the kind of soil at research location was andosol and regosol. The average of research location altitude was 2100 m from sea level. According to previous research, kidney bean good growth on altitude (2,000-2,500) meters from sea level and the kind of soil was regosol and andosol (Rukmana, 2003). Characteristics of andosol soil were black color, clay to dust texture, loose and medium permeability (Rukmana, 2003). Characteristics of regosol soil were gray and yellow color, send texture and permeable.

The variation of temperature at the research location was 18-28°C on the planting climate. The average temperature of five research locations was 24°C. Previous research reports that the ranges temperature for good growth of kidney bean was 20-25°C (Rukmana, 2003).

The average rainfall at the research location was 1,650 mm/year. According to previous research, kidney bean can good growth on rainfall between (1,500-2,500) mm/year (Rukmana, 2003). The variation of humidity at the research location was (48-64%). This reason, on humidity between (70-80%), the several aphids can be more past breed.

Based on data in Table 1, 2 and 4: Data calculations, observation and explanation above can be suggested any arguments that mutant that more quickly flowered, more fertile, mass per 1,000 seeds is higher, growth rate percentage is higher than control, which was adapted to germ, dry condition, high calcium, high salt, the mean or average production is higher. The mean production of control sample (initial variety) is 2.60 tons/ha and mean production of mutant (generation variety) is 4.71 tons/ha.

This result shows that mean production of mutant of local kidney bean from Manggarai Flores Indonesia was increased significantly. The increase of mean or average production is 44.80%.

Conclusion

Based on explanation upon can be proposed that development of local kidney bean from Manggarai Flores Indonesia by irradiation with using multi gamma radiation source and carefully selection was obtained mutant seeds of kidney bean with physical and chemical characteristics was superior for continuously development. The range production of mutant was obtained from mutation by irradiation with using multi gamma radiation was 3.97 tons/ha up to 5.28 tons/ha, mean or average production was 4.71 tons/ha and the
increase of mean production was 44.80%. The treatment samples selected were tolerant to area with dry conditions, high calcium and salt and tenacity to germ.

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Author’s Contributions

The all authors have contributions of this manuscript like as collaborated on research and always to be discussed for writing manuscript.

Conflict of Interest

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