

# Physiological and genetic aspects of a diploid potato population in the Netherlands and Northern Finland

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## Abstract

Tuberization of potatoes exposed to different photoperiod regimes has been earlier investigated in several studies. However, there is still a limited understanding of the entire tuberization process and the factors influencing this process. One of the constraints of the previous studies has been the use of only one or a few genotypes. Furthermore, the experimental designs have not utilised the natural growing conditions with continuous changes in day-length during the growing season. The general aim of the project was to study the developmental dynamics of the broad-based potato (*Solanum tuberosum* L.) (CxE) population development at different climatical regimes under the very long-day, long-day and short conditions in Finland, the Netherlands and Ecuador/Venezuela, respectively. In this paper we are presenting some of the results achieved in the studies in Finland during the growing season 2004. In addition to population level trait characteristics we also describe here some of the identified QTLs (quantitative trait loci) for stolon related and tuber formation traits. In some cases we also compare the expression of some of the traits both in Finland and Netherlands.

The main difference between the CxE population grown in Finland and the Netherlands was that the onset of flowering took place one week later in Finland. The relationship between tuber initiation and onset of flowering differed depending on the day length. In Finland approximately 70 % of the genotypes had swollen stolon tips before the onset of flowering, while in the Netherlands only 30 % of the genotypes had reached this condition. We also found numerous different trait linked QTLs, for example, a QTL associated with tuber formation was identified on chromosome E5, and QTLs associated with stolon characteristics on chromosomes E1, E4, E10 and E12. As a conclusion, the present preliminary results provide a good basis for determining the influence of different environmental conditions on potato development. In addition, the QTLs obtained in this study give a better understanding of the genetics of complex characters, and can be used in improving the potato crop in breeding programs.

## Key words

Potato, flowering, tuberization, day length, QTLs

## Introduction

Tuberization of potatoes exposed to different photoperiod regimes has been earlier investigated in several studies (Werner 1942, Driwer & Hawkes 1943, Mendoza & Haynes 1976, Ewing 1978, Regel & Sands 1983, Khedher & Ewing 1985, Martinez-Garcia et al 2002). However, there is still a limited understanding of the entire tuberization process and factors influencing this process. One of the constraints of the previous studies has been the use of only one or a few genotypes. Furthermore, the experimental designs have not utilised the natural growing conditions with continuous changes in day-length during the growing season.

The general aim of the project is to compare the dynamics of the genetically broad based potato population development in Finland, the Netherlands and Ecuador / Venezuela in order to gain understanding on how different environmental conditions, especially day length, affect expression of various traits. In this paper we are presenting some of the results achieved in the studies in Finland during the growing season 2004. In addition to population level trait characteristics we also describe here some of the identified QTLs (quantitative trait loci) for stolon related and tuber formation traits. The QTL analyses were done in order to gain more knowledge on the genetics of these traits, to examine their expression at different developmental stages, and to investigate the potential environmental impact. The analyses were realised due to the existing molecular marker data produced at the Laboratory of Plant Breeding in Wageningen University. In some cases we were also able to compare the expression of some of the traits both in Finland and Netherlands.

## Material and methods

The diploid backcross population termed “CxE” with 232 genotypes and 20 tetraploid cultivars which were used as controls, which are typically grown either in the Netherlands or in Finland, were supplied by the Laboratory of Plant Breeding, Wageningen University for the field experiments. The parental clone C is a hybrid between *S. phureja* PI225696.1 and *S. tuberosum* dihaploid USW42, while the parental clone E is the result of a cross between clone C and the *S. vernei*-*S. tuberosum* backcross clone VH<sup>3</sup>4211 (Celis-Gamboa et al 2002, 2003, Celis Gamboa 2002).

In this paper we are presenting mainly results from the field experiment conducted at the North Ostrobothnia Research Station in Ruukki (64°42' N, 25°00' E) in 2004, but also from the previous corresponding experiment carried out in the Netherlands (51°58' N, 5°38' E) in 1999. AFLP markers and map, which were generated and constructed by Celis-Gamboa (2002) on the CxE population, were used in the QTL analyses.

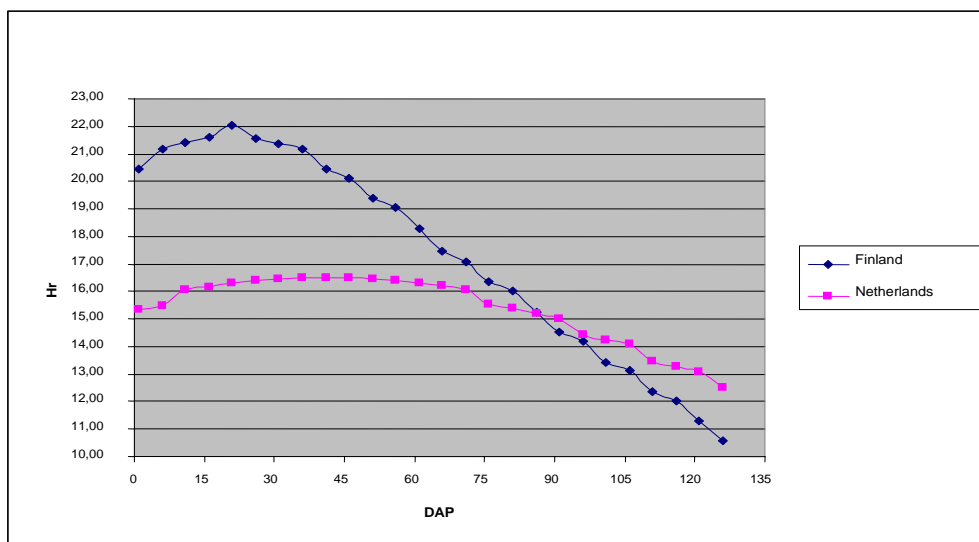


Figure 1. Differences in day length (hours of light) during the growing season in Finland and in the Netherlands. The day length is presented by days after planting (DAP).

The experimental design for field experiment in Finland followed the one in the Netherlands. Each genotype with 3 individuals from the CxE population was randomised within 8 blocks. The blocks from 1 to 7 were harvested with start on 30 days after planting (DAP) in approximately 7 days intervals. Block 8 was harvested in four steps from 112 DAP after the senescence of the individual genotypes. Each of the blocks included also 20 control varieties, with 3 individuals (altogether 60 control plants). The experiments were managed by using standard potato cultivation methods of the respective experimental site. The most important environmental factor differing between the experimental sites was the day length during the growing season (Figure 1.). All the evaluations were conducted on all the three individuals of each genotype and the control varieties. Altogether, eighteen developmental traits were measured (Table 1).

Table 1. Developmental traits of the (CxE) potato population evaluated in the very long-day field trial experiment in Finland in 2004. Evaluated characteristics present both above and below ground traits.

Evaluated trait	Description	Evaluated trait	Description
1. Date of emergence	The days from planting to emergence (DAP)	10. Degree of stolon branching (scale)	Scale 1-5 (1 = no branching, 5 = more than 20 branches)
2. Number of main stems	Stems emerging directly from the seed tuber	11. Number of swollen stolon tips (stolon forming tubers)	A tuber was defined as swollen tip with at least 12 mm in diameter.
3. Number of wild stems	Stems with leaves that started as stolons followed by a switch to orthotropic growth	12. Position of swollen stolon tips and growing tubers	Stolon tip, on a stolon side branch, as part of a chain on the stolon, or other locations
4. Plant height (cm)	Length of the highest stem from the ground level to the main apex	13. Secondary tuber growth (scale)	Scale 1-5
5. Flower colour (for control of the genotype)	White, red violet, blue violet, light blue or other possible colours	14. Tuber size distribution (mm)	Each tuber larger than 12 mm in diameter was recorded.
6. Flowering (scale)	Scale from 1-7 (1 = first open flower, 4 = peak of flowering, 7 = end of flowering).	15. Total tuber weight per plant (g)	
7. Senescence (scale)	Scale 1 to 7 (1 = first yellow leaves, 7 = no green tissue left)	16. Quality of root system (scale)	1 = poor, 2 = normal, 3 = very long and dense
8. Number of new stolons	Count of all the stolons that do not show signs of tip swelling	17. Length of root system (cm)	
9. Length of the longest stolon (cm)		18. Incidence of <i>Streptomyces scabies</i> and <i>Rhizoctonia solani</i>	Visual inspection of the harvested tubers

The statistical analyses were carried out with the SPSS 12.0 program. The data was tested for outliers, and if found, they were excluded from the calculations. All the traits were tested for normality and when needed transformed (square root or log<sub>10</sub> transformation). Descriptive statistics e.g. means, standard deviation, minimum and maximum of the CE population, for different traits were calculated. Bivariate 2-tailed Pearson-Correlation was used to calculate correlation between different traits both for the Finnish and Dutch data. When needed the data from the Netherlands was also transformed (log transformation) to fit the data from Finland in the correlation test.

For generation of AFLP markers 15 *EcoRI/MseI* based primer combinations were used, and the markers were visually scored as presence or absence of the band. The linkage analysis was performed according to the double pseudo-testcross approach for cross pollinating populations and using the

software JoinMap 2.0. The 403 markers obtained from the AFLP analysis, were used to construct a maternal (C) and a paternal (E) maps with 12 linkage groups each. The C map consisted of 140 markers spanning 919 cM and the E map consisted of 178 markers spanning 784 cM. In order to identify if the origin of the map each linkage group is preceded by the letter C or E, followed by the linkage group number (Celis-Gamboa 2002).

For the QTL analysis the data was evaluated at first for normality with normal Q-Q plot-proportion estimation formula according to Blom (1958). Rank assigned to ties: Mean, which showed the correlation between the normal expected values to the observed values, and when needed the data was transformed. Differences between genotypes were evaluated using univariate analysis, and the mean values for each genotype obtained from the univariate test were used in the QTL analysis. Since the maternal (C) and the paternal (E) maps could not be integrated unambiguously, due to the low informative value of the dominantly scored 3:1 AFLP “bridge” markers, QTL mapping was performed on the separated C and E parental maps.

Using the software package MapQTL version 5.0 (Van Ooijen, 2004), the mean values of normally distributed quantitative traits were first analysed by interval mapping. The program was set to calculate in steps of 5cM between the linkage markers and the criteria for detecting QTL with significance level of 0.05, was defined for threshold LOD value of 3.0, after performing a permutation test, as implemented in MapQTL version 5.0. In the next step cofactors were tested according to the results obtained from the interval mapping. Using the automatic cofactor selection options, all the markers with LOD values larger or close to 3.0, were tested. The selected cofactors were then used to run the multiple-QTL model test, which gave rise to more accurate peak locations. To obtain 95% confidence interval around the estimated point, a so called two-LOD support interval was constructed, by taking the two positions, left and right to the peak, that had a LOD value of two less than the maximum. All the identified QTLs, were compared to the QTLs identified in the CxE population grown in the Netherlands.

## Results and discussion

### *The dynamics of the CxE population development in Finland*

Highly significant differences ( $P < 0,01$ ) were found between the genotypes for the traits evaluated. In Figure 2. developmental patterns of the CxE population in Finland are demonstrated.

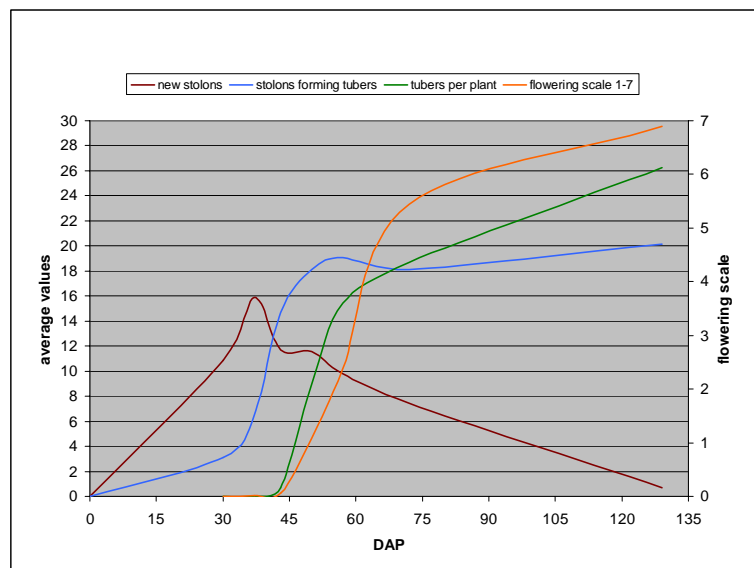


Figure 2. A general view of the development of the CxE population in Finland.

The main features of the developmental processes on the population level were the formation of stolons and tuber incipients, as well as the start of stolon branching, well before start of flowering. In the corresponding experiment in the Netherlands these processes occurred somewhat later in relation to flowering (Celis-Gamboa et al. 2002). Initiation of stolon swelling (10 days difference) and

emergence of first tuber incipient (7 days) in the varieties took place on average 10 days earlier than in the CxE clones (Zaban 2005).

#### ***Association between flowering and tuberization***

In the Netherlands 30% of the CE population had swollen stolon tips before, 34% simultaneously with and 36% after the first flower was opened. In Finland the corresponding values were 71, 21 and 8 % (Figure 3). In the Netherlands most of the CE population started flowering between 36 and 42 DAP, and the mean for onset of flowering for all the varieties was on 42 DAP, while in Finland the mean onset for flowering for CxE population was 50 DAP. No correlation was found between the onset of flowering (DAP) in Finland and the onset of flowering (DAP) in the Netherlands ( $r = -0,039$ ). These results indicate contrary to earlier studies (reviewed by O'Brien et al. 1998) that the initiation of stolon tip swelling is not dependent on the onset of flowering. However, no conclusions on the relationship between tuber induction and flowering can be drawn from our data. More information on this aspect has been gathered in the field experiment of the growing season 2005.

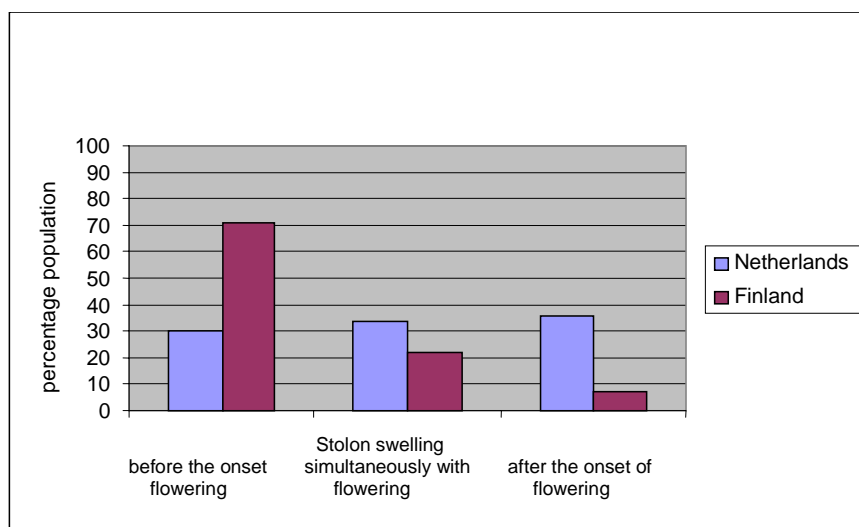


Figure 3. Relationship between flowering and tuber initiation in Finland and the Netherlands.

#### ***QTLs***

As a result of QTL analyses a QTL associated with tuber formation was identified on chromosome E5, and QTLs associated with stolon characteristics were identified on chromosomes E1, E4, E10 and E12 (Table 2).

All the QTLs identified both in Finland and in the Netherlands were compared to examine if they share similar locations. Figure 4 illustrates schematically QTLs detected in the chromosome e5. The results may indicate that the traits are, at least partially, interrelated and are controlled by the same genes (Simko et al. 1999). The same QTL as identified for the number of stolon forming tubers at 37 and 43 DAP on chromosome e5, was found in the Netherlands associated (with very high LOD values) to the duration of plant life cycle, onset of senescence, mid point of senescence and plant maturity type (the population was classified in categories according to the duration of the plant life cycle) (Celis-Gamboa 2002). Another trait from the Netherlands, which had the same QTL as the trait for number of stolon forming tubers at 37 and 43 DAP, was the number of tubers per plant. The QTL identified in Finland to the number of stolon forming tubers at 50 DAP, was associated in the Netherlands to the duration of plant cycle and the duration of senescence. Also the QTL results on tuber formation obtained in this study are in agreement with earlier studies (Van den Berg et al. 1996, Schäfer-Pregl et al. 1998, Simko et al. 1999), a fact that confirms the relevancy of the data collected.

Table 2. QTLs identified for the degree of stolon branching and for the length of the longest stolon in the field trial in Finland.

Trait	QTLs information			
	Chrom.	Position	LOD	Marker
Stolon branching 50 DAP	e4	53.1	4.52	E39/M60-42e4
Stolon branching 72 to 129 DAP	e4	52.1	9.21	E32M50-206e4
	e10	75.5	3.12	E32M54-74e10
	e12	51.7	3.49	E39/M60-9e12
Stolon length 50 DAP	e1	16.4	4.24	E38/M59-431e13
		30.1	5.7	E32M61-18e13
	e4	52.1	4.82	E32M50-206e4
Stolon length 72 to 129 DAP	e1	16.4	4.02	E38/M59-431e13
		30.1	4.83	E32M61-18e13
	e4	54.6	9.4	E32M50-105e4
	e12	59.2	3.09	E32M50-97e12

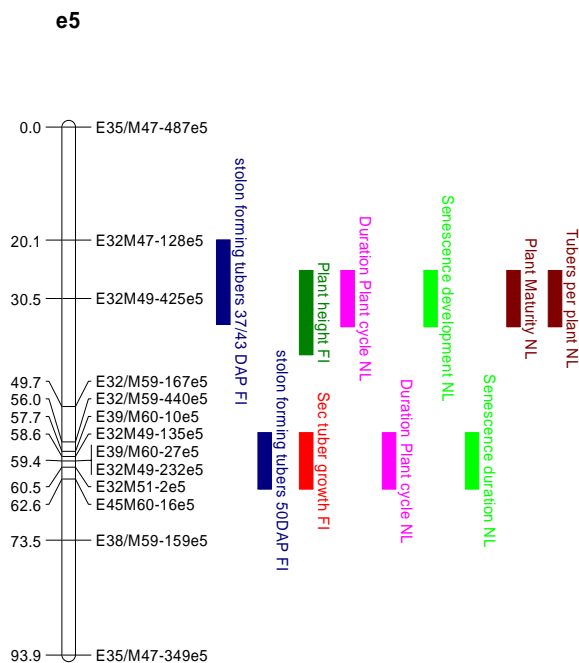


Figure 4. Schematic overview of QTLs detected on chromosome e5 in Finland and the Netherlands. Indicated are the QTL peak positions and an arbitrary QTL support interval of 5 cM on both sides of the peak position.

## Conclusions

It is well known that repeated QTL mapping studies, using identical materials in different environments or years will give varying results. This is due to the limited power of detection of QTLs in practically manageable population sizes, which, together with random experimental effects can cause a different subset of the genetic effects present in the population to manifest itself in each single analysis, especially for smaller effect factors that just surpass the detection limit. This is why repetition of experiments is essential to identify true factors that underlie traits of interest from one-time analysis artefacts.

As a conclusion, this is the first time that the effects of environmental conditions and especially different day length on potato development can be followed and compared. This work has been possible due to the use of the same potato population, molecular marker map and field trials established in the Netherlands and in Finland. The preliminary results obtained in this study (mainly from Finland from one growing season 2004) and the already existing data from the experiment in the Netherlands provide a good basis for determining the influence of different environmental conditions on potato development. More accurate analyses will be realised when the data collected from the follow-up field experiment in Finland 2005 has been analysed. Already the preliminary data achieved indicates interesting associations between tuber initiation and onset of flowering and some common, thus meaningful, QTLs in data from obtained in Finland and the Netherlands. Further QTL analyses including those from S America will provide better understanding of the genetics of complex characters during potato development and, furthermore, will lead to the improvement of the potato crop via breeding programs.

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