

E. coli Genome Manipulation by P1 Transduction

Lynn C. Thomason,¹ Nina Costantino,¹ and Donald L. Court¹

¹National Cancer Institute at Frederick, Frederick, Maryland

ABSTRACT

This unit describes the procedure used to move portions of the *E. coli* genome from one genetic variant to another. Fragments of ~100 kb can be transferred by the P1 bacteriophage. The phage is first grown on a strain containing the elements to be moved, and the resulting phage lysate is used to infect a second recipient strain. The lysate will contain bacterial DNA as well as phage DNA, and genetic recombination, catalyzed by enzymes of the recipient strain, will incorporate the bacterial fragments into the recipient chromosome. *Curr. Protoc. Mol. Biol.* 79:1.17.1-1.17.8. © 2007 by John Wiley & Sons, Inc.

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INTRODUCTION

P1 transduction is a useful genetic procedure for moving selectable mutations of interest from one *E. coli* strain to another. P1 is a bacteriophage with a rather sloppy packaging mechanism, and during growth and encapsidation of its DNA into the phage head, it occasionally packages the DNA of its bacterial host rather than its own phage chromosome into the protein capsid. This property of the P1 phage means that when a P1 lysate is made, the individual particles in the lysate contain either packaged phage DNA or packaged bacterial DNA. When this lysate is used to infect a second (recipient) host, DNA pieces from the original host chromosome are transferred into the new strain, where they can recombine and be permanently incorporated into the chromosome of the second strain, an event called transduction. Successful P1 transduction of any one marker from a unique location in one bacterial strain to a second strain is a relatively rare event, so selection is required. P1 can carry two or more different genetic markers in the same transducing particle DNA, and at some frequency both of these markers can be recombined into the recipient strain. Such markers are said to be linked by P1 transduction, i.e., they are close enough to be co-transduced. Proximity of the selectable marker to the allele of interest is important, because nearer alleles will co-transduce more frequently. The allele of interest might be a (defective) gene with an antibiotic marker cassette insertion, or it might be a gene with a point mutation that is closely linked to a selectable antibiotic marker.

To execute the Basic Protocol, the P1 lysate is grown on a host containing a genetic variant of interest and this lysate is then used to infect a second strain. For the nonexpert, it is best to use a virulent phage mutant, P1 *vir*, to produce the lysate because the *vir* mutation prevents the phage from generating P1 lysogens among transductants. The Basic Protocol describes growth of a liquid lysate, which is easier to carry out than a plate lysate and generally gives good results. The Support Protocol describes the procedure used to titer the P1 stock, which is usually done only if the transduction is not successful. The Alternate Protocol provides a method for preparing a plate lysate, which may be used if the starting P1 preparation has a low titer. Plate lysates often yield higher-titer stocks than liquid lysates.

PREPARATION OF P1 LIQUID LYSATE AND P1 TRANSDUCTION

This protocol describes the two-part procedure: preparation of a liquid P1 lysate, followed by transduction. Each procedure requires a freshly grown bacterial overnight culture. Once the cultures are prepared, the procedures are simple and each requires only a few hours, so that both can be performed in the same day.

Materials

- LB broth (*UNIT 1.1*)
- E. coli* nonlysogenic donor strain (containing the genetic marker to be transduced)
- 20% glucose stock solution
- 1 M CaCl₂
- 10⁹ to 10¹⁰ pfu/ml P1 stock, a virulent P1 mutant preferred (e.g., P1_{vir})
- Chloroform (CHCl₃)
- E. coli* recipient strain (to be transduced)
- Tris-magnesium (TM) buffer: 10 mM MgSO₄/10 mM Tris·Cl, pH 7.4 (*APPENDIX 2*)
- P1 salts solution: 10 mM CaCl₂/5 mM MgSO₄
- 1 M sodium citrate
- Selective plates (see *UNIT 1.1*) with 5 mM sodium citrate (recommended); e.g., minimal plates if selecting for prototrophy *or* rich LB plates containing the appropriate antibiotic (depending on drug cassette used):
 - 30 µg/ml ampicillin
 - 30 µg/ml kanamycin
 - 10 µg/ml chloramphenicol
 - 12.5 µg/ml tetracycline (6.25 µg/ml if sodium citrate is added)
 - 50 µg/ml spectinomycin
- 50-ml Erlenmeyer flasks, preferably baffled
- 18 × 150-mm glass culture tubes
- Shaking water bath or test-tube roller at appropriate temperature (usually 32° to 37°C)
- 35-ml plastic centrifuge tubes
- Refrigerated low-speed centrifuge with Sorvall SA-600 rotor (or equivalent)
- 5-ml screw-cap vials

NOTE: All material coming in contact with *E. coli* must be sterile.

Prepare liquid P1 lysate

1. Inoculate 5 ml LB broth with the *E. coli* donor strain (from which the P1 lysate is to be made). Incubate overnight at the appropriate temperature.

Throughout this protocol, the optimal temperature for cell growth is 37°C, unless a temperature-sensitive strain is used. Temperature-sensitive strains should be cultured at 30° to 32°C.

To perform transduction on the same day, grow the recipient strain (step 9) at the same time.

2. Dilute the overnight donor culture 1:100 into 5 ml LB broth containing 50 µl 20% glucose (final concentration 0.2%) and 25 µl of 1 M CaCl₂ (final concentration 5 mM) in a 50-ml Erlenmeyer flask or 18 × 150-mm glass culture tube.
3. Incubate 30 to 45 min at 30° to 37°C with shaking.

Use either a 50-ml Erlenmeyer flask in a shaking water bath or a standard culture tube with adequate aeration in a test-tube roller. If the strain is temperature sensitive, use the lower temperature and the longer time.

4. Add 100 μl of a recently prepared 10^9 to 10^{10} pfu/ml P1 vir stock.

A good titer for a P1 stock is 10^9 to 10^{10} pfu/ml. If the stock has recently been used successfully and does not contain CHCl_3 (which reduces the viability of the stock; see step 6), it is probably unnecessary to determine the titer.

5. Continue to incubate, with shaking, until the culture lyses (~ 3 hr).

The culture may not lyse and clear completely, but clumps of floating debris should be visible when it is held up to the light. An unlysed culture looks smooth and silky when swirled; a lysed culture does not.

6. Add a few drops of CHCl_3 and continue shaking a few minutes more.

The CHCl_3 ensures complete cell lysis and kills the bacteria. CHCl_3 does not harm the P1 phage at this step, but care should be taken to avoid transferring the CHCl_3 to the storage tube during step 8, because storing the phage in the presence of CHCl_3 can result in decreased viability of the stock over time.

7. Transfer the supernatant to a 35-ml centrifuge tube and centrifuge 10 min at $\sim 9200 \times g$, 4°C .

8. Pour off the supernatant (the P1 lysate) into a screw-cap 5-ml tube. Label it with the strain on which the lysate was grown (e.g., P1 vir·W3110) and the date, and store up to several years at 4°C .

The lysate can be further purified by passage through a sterile 0.45- μm filter syringe after centrifugation. The filter will remove any residual bacteria.

The lysate can be stored for several years at 4°C and still retain adequate titer. If in doubt, either spot titer as described in the Support Protocol or make a new lysate.

Perform P1 transduction

9. Inoculate 5 ml LB broth with the *E. coli* recipient strain. Incubate overnight at the appropriate temperature.

If the strain does not have temperature-sensitive elements, use 37°C .

10. The following day, centrifuge 1.5 ml of the cell culture in a microcentrifuge tube 2 min at maximum speed, room temperature, to pellet the cells.

11. Discard the medium and resuspend the cells at one-half the original culture volume in sterile P1 salts solution.

This gives 0.75 ml suspended cells, which is adequate for two transductions. Process more cells in step 10, if necessary.

12. Mix 100 μl of the cells/P1 salts mixture with varying amounts of lysate (1, 10, and 100 μl) in sterile test tubes. As a control, include a tube containing 100 μl of cells without P1 lysate.

13. Allow phage to adsorb to the cells for 30 min, either at room temperature on the benchtop or at the appropriate culture temperature. Treat the controls similarly.

Temperature selection should take into account any temperature or cold sensitivity of the recipient strain.

14. Add 1 ml LB broth plus 200 μl of 1 M sodium citrate.

The sodium citrate chelates the calcium essential for phage adsorption to the bacteria, thus minimizing secondary infection by residual phage.

15. Incubate ~ 1 hr at 30° to 37°C with aeration.

The exact temperature will depend on the characteristics of the strain being transduced and the expected characteristics of the strain after transduction. If the desired strain is temperature sensitive, the incubation should be done at a low temperature.

16. Centrifuge each culture in a 1.5-ml microcentrifuge tube 2 min at maximum speed, room temperature, to pellet the cells. Remove and discard the supernatant.
17. Suspend the cells in 50 to 100 μ l of an appropriate buffer or broth and spread the entire suspension onto the appropriate selective plate. Also spread 100 μ l of the P1 stock on another plate. Incubate the plates overnight at the appropriate culture temperature.

It is good policy to supplement the selective plates with 5 mM sodium citrate, which helps prevent readsorption of residual P1 phage. If sodium citrate is used, one-half the concentration of tetracycline must be used.

Antibiotic resistance is a commonly selected marker: in this case, LB broth can be used for dilution. Be sure to use plates with the lower drug concentrations needed for single-copy genes. Prototrophy can also be selected: in this case, the cells should be washed and diluted in a minimal salts medium such as TM buffer.

The P1-only control is used to determine whether contaminating bacteria are present in the lysate.

18. The next day score the plates.

The control plates with cells only or P1 phage only should be free of bacterial colonies; the other plates should contain the transduced colonies. Usually the plate receiving the lowest amount of P1 lysate will have the fewest colonies.

If colonies are present on the cells-only control plate, either the selection is not working or contamination is present.

If colonies are present on the phage-only plate, the P1 lysate is likely contaminated with donor bacteria. This should not occur if the CHCl_3 treatment was complete. If this control gives colonies, filter the lysate through a sterile 0.45- μ m filter and try again.

19. From the plate that shows growth from the least amount of P1 phage, pick single colonies, streak to isolate single clones, and incubate plates overnight at 30° to 37°C. Repeat this process a second time.

Picking colonies from the plate to which the least amount of P1 was added is good microbial practice because many infective P1 phage particles will remain on the plates. Two rounds of purification are essential to ensure removal of such residual P1 phage and separation of nontransduced bacteria from the actual transductants.

Table 1.17.1 Correlation of Linkage with Physical Distance for P1 Transduction

Percentage linkage	Physical distance (kb)
0	—
2	67.03
10	49.30
20	38.20
30	30.41
40	24.21
50	18.98
60	14.40
70	10.31
80	6.59
90	3.17
100	0.00

If the genetic marker selected is identical to the mutation of interest (100% linkage), every colony should express the mutant phenotype. If the genetic marker selected is not tightly linked to the mutation, more colonies must be screened for the desired phenotype (see Table 1.17.1).

20. Confirm the genotype of the purified transductants by drug-resistance or phenotype.

If there is no genetic way to score the desired mutation, use PCR to amplify the region of interest (using standard primers) and sequence the PCR product.

TITRATION OF THE P1 LYSATE

Although P1 transduction is usually performed without titering the phage lysate, the phage should be assayed if the transduction is not successful. This protocol describes spot titering, where the lysate is serially diluted and applied to a bacterial lawn. Spot titering conserves materials and may facilitate visualization of the P1 plaques, which are very tiny.

Additional materials (also see Basic Protocol)

Standard *E. coli* K12 strain (e.g., C600 or MG1655)
LB plates and top agar (UNIT 1.1), each containing 2.5 mM CaCl₂

1. Inoculate 5 ml LB broth with a standard *E. coli* K12 strain. Incubate overnight until cells reach mid-log phase ($\sim 2 \times 10^8$ cells/ml).
2. Add 1 M CaCl₂ to a final concentration of 5 mM.
3. While the cells are growing, make serial ten-fold dilutions (through 10^{-8}) of the phage stock in TM buffer.
4. Pour a lawn of the cells on an LB/CaCl₂ plate by mixing 0.25 ml of the cell culture with 2.5 ml LB top agar (with CaCl₂) in a small culture tube and immediately pouring the contents onto the plate. Tilt the plate gently to distribute the cell-agar mixture uniformly across the surface of the plate.
5. After the agar hardens, spot 10 μ l of each phage stock dilution onto the lawn of cells. Let the spots dry and then incubate the plate upright at 37°C.

After overnight incubation, individual countable plaques should be present in one of the dilutions.

6. Calculate the titer of infective phage particles/ml (pfu/ml) by counting the number of plaques and multiplying by the fold dilution and a factor of 100 (to account for the 10 μ l volume spotted).

If the only P1 lysate available has a poor titer ($< 10^9$ pfu/ml), a new high-titer stock must be grown. In this case, use a confluent 10- μ l spot of phage to inoculate a liquid culture and make a high-titer liquid lysate (see Basic Protocol, steps 1 to 8). The spot can be taken from the lawn of cells using a sterile spatula (do not dig into the bottom agar).

PREPARATION OF P1 PLATE LYSATES

Plate stocks can give higher titers than liquid stocks, although they are often not necessary. This is a standard procedure for making P1 plate lysates. Variations exist and may also work well. For materials, see Basic Protocol.

1. Inoculate 5 ml LB broth with the *E. coli* donor strain. Incubate overnight at the appropriate temperature.

The optimal temperature is 37°C, unless a temperature-sensitive strain is used. Temperature-sensitive strains should be cultured at 30° to 32°C.

SUPPORT PROTOCOL

ALTERNATE PROTOCOL

*Escherichia coli,
Plasmids, and
Bacteriophages*

1.17.5

2. Dilute the overnight donor culture by inoculating 100 μ l into 5 ml LB broth containing 0.1% glucose. Incubate \sim 1.5 hr until cells reach mid-log phase ($\sim 2 \times 10^8$ /ml).

Use either a 50-ml Erlenmeyer flask in a shaking water bath or a standard culture tube with adequate aeration in a test-tube roller.

3. Add 1 M CaCl_2 to a final concentration of 5 mM.
4. In a standard 18 \times 150-mm glass culture tube, mix 3 ml culture with 50 μ l of a 10^9 to 10^{10} pfu/ml P1 *vir* stock.
5. Incubate the phage/cell mixture 20 min at 37°C or room temperature for the phage to adsorb.
6. Add 7.5 ml melted LB top agar to the phage/cell mix and distribute it among three fresh LB plates supplemented with 0.1% glucose and 2.5 mM CaCl_2 . As a control, pour a plate containing cells alone. Incubate the plates upright overnight.

Do not invert the plates because the top agar is very soft.

Following incubation, the bacterial lawn will appear lysed and may contain mucoid colonies. Compare the appearance of the control lawn to the plate lysates. A smooth bacterial lawn is an indication that the phage lysate did not grow well.

7. To harvest lysates, scrape the top agar from each plate into a 35-ml centrifuge tube. Wash each plate with 2 ml LB broth and add to the tubes.
8. Add a few drops of CHCl_3 to each tube and vortex hard for 30 sec. Let the tubes stand on the bench for 10 min.
9. Centrifuge the tubes 10 min at $\sim 14,500 \times g$, 4°C.
10. Decant the supernatant into a labeled screw-cap tube and store the lysate up to several years at 4°C.

The lysate can be stored for several years at 4°C and still retain adequate titer. If in doubt, either spot titer as described in the Support Protocol or just make a new lysate.

COMMENTARY

Background Information

The use of P1 as a transducing phage was first reported in the early days of molecular biology (Lennox, 1955), and it has become a standard tool for microbial genome manipulation. P1 is a generalized transducing phage, meaning that it will package, and is thus capable of transducing, the entire *E. coli* genome, although it may have preferred sites in the bacterial DNA where packaging is initiated. After initiation of packaging somewhere on the bacterial chromosome, the DNA is encapsidated processively by a headful mechanism. Transducing particles conservatively make up \sim 0.1% of the total number of phage particles in a lysate (Yarmolinsky and Sternberg, 1988), and \sim 0.1% to 1% of these will contain the particular DNA one wants to move into another bacterial strain. The capacity of the P1 head is \sim 100 kb; thus \sim 100 kb of contiguous bacterial DNA can be moved from one bacterial strain to another.

Other P1 phage types have been used for transduction, for example: P1*kc*, which forms clear plaques (Lennox, 1955); P1Tn9*clr*100, a P1 derivative with a chloramphenicol resistance gene and a temperature-sensitive repressor (Silhavy et al., 1984); and hyper-transducing variants (created by Nat Sternberg), which package the bacterial DNA more frequently (D. Friedman, pers. comm.). However, all these variants can generate serious problems because they are capable of lysogeny. In contrast to phage λ , phage P1 lysogens maintain the phage chromosome as a plasmid rather than integrated into the bacterial chromosome.

P1 transduction is especially useful in conjunction with the recently developed technique of recombinering (UNIT 1.16), a method for in vivo genetic engineering in *E. coli*. The technique of recombinering allows creation of nearly any desired mutation or genetic construct. Providing there is a selection method

for the construct, a genetic construct can be created by recombineering and then moved into a different genetic background by P1 transduction.

Critical Parameters and Troubleshooting

Microbiological methodology

P1 transduction is a genetic procedure requiring sterile microbiological methods. Glassware and solutions must be sterile, and care should be used throughout this procedure to avoid contamination. Transductants must be purified selectively two consecutive times, preferably on plates containing sodium citrate, to ensure elimination of residual P1 phage. The citrate chelates calcium important for efficient phage adsorption to the bacteria, thus minimizing infection by the remaining phage.

P1 stock

A reasonable P1 stock titer is 10^9 to 10^{10} plaque-forming units (pfu)/ml. A fresh stock should be obtained from a known source before beginning this procedure. It is good practice to use a P1 stock grown on a wild-type *E. coli* strain to ensure a high-titer lysate. However, any P1 lysate of good titer can be used to generate a new lysate for transduction, because bacterial DNA present in the first lysate is not replicated and will not contaminate the second lysate. Most investigators rarely titer P1 stocks; however, if the transduction is not successful, the phage stock should be titered because a low titer will prevent isolation of transduced colonies. Once made, the stock will probably be good for at least a year if it is not stored over CHCl_3 , and older stocks can be used if they maintain titer. Storage in the presence of CHCl_3 will cause the titers to drop appreciably within a few months.

Linkage

The closeness of two markers is indicated by the frequency (as a percentage) of their co-inheritance (see Table 1.17.1). Often the allele to be transduced is linked to an antibiotic resistance gene, and drug resistance is selected. Prototrophy for a metabolic gene can also be selected, e.g., the ability to grow in the absence of an amino acid. In both cases, the presence of the second allele must later be confirmed by phenotype or sequencing. Recombineering (*UNIT 1.16*) can be used to insert a selectable marker near a gene or mutation that cannot readily be selected; P1 transduction can then be used to move the drug marker and the linked mutation into a different strain. The laboratory

of Carol Gross has created a set of bacterial strains with drug insertions at about 1-min intervals around the *E. coli* chromosome (Singer et al., 1989). A collection of *E. coli* single-gene knockout mutants (the Keio collection) has also recently been described (Baba et al., 2006). P1 can be grown on any of these strains and transduction used to move the drug marker in a particular region of the genome into another more desirable background.

Host elements that interfere with transduction

Once a recipient strain has been infected with the P1 lysate, incorporation of the bacterial DNA into the recipient chromosome requires host recombination functions, especially RecA and RecBCD. It is thus not possible to transduce a *recA* mutant strain, since absence of the RecA function reduces homologous recombination ~ 1000 -fold. Mutations in the RecBCD enzyme also affect this procedure adversely, although these mutants are less defective for recombination than are *recA* mutants. To use P1 transduction to build an *E. coli* strain with multiple mutations, including one in *recA*, the *recA* mutation should be incorporated last. These recombination-defective strains are also poor hosts for phage P1 and give lower-titer lysates. Mutations in other recombination functions may also have a detrimental effect. At least one non-recombination-related mutation also impacts P1 growth. Phage P1 adsorbs to a sugar residue of the lipopolysaccharide (LPS) on the *E. coli* cell surface (Sandulache et al., 1984), and a mutation in the *galU* gene, which encodes a component of the LPS, makes *E. coli* unable to support the growth of phage P1 (Franklin, 1969).

Be extremely careful of using any P1 stock made on bacterial lysogens containing phages such as λ or Phi80. These other phage types will be present in the P1 stock and can cause serious contamination problems when successive P1 stocks are made from them. An exception is the defective λ prophage present in the recombineering strains referred to in *UNIT 1.16*. Since this prophage is incapable of excision from the bacterial chromosome, it cannot contribute contaminating λ phage particles.

Anticipated Results

Since one in $\sim 10^3$ phages in a P1 lysate are transducing phage, and phage P1 can package 1% of the *E. coli* chromosome, each 1% of the bacterial chromosome will be represented by one transducing phage in a population of 10^5 total phage. Since a high-titer lysate contains

~10⁹ phages/ml, and since the frequency of incorporation into the recipient by recombination will be <100%, for any one marker typically there will be ~100 to 1000 successful transduction events/ml of lysate. This rough calculation assumes equal transduction of all parts of the *E. coli* chromosome, although in reality some regions of the genome are “hotter” and will transduce better than others.

Time Considerations

The entire procedure of growing the overnight cultures, making the P1 lysate, performing the transduction, and purifying the colonies will take less than a week. Once the lysate is grown and the transduction is complete, the steps on subsequent days are simple and should only take a few minutes of actual hands-on time.

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These are classical references for transduction by phage P1 and other phages.

Internet Resources

<http://rothlab.ucdavis.edu/protocols/p1-transduction-chart.html>

This Web site contains a more complete version of Table 1.17.1.