

Interprocedural Dataflow Analysis

Propagating information across procedure boundaries is useful.

- Optimize caller using information about callee

```
x := 2
```

```
CALL f(x) //call by reference
```

```
...x... //is x equal to 2 here?? Need to look at f
```

- Optimize callee using information about callers

```
... PROCEDURE f(a,b)
```

```
CALL f(x,5) .....
```

```
...
```

- We might generate specialized code for f in which b is 5.
- We might *clone* f and specialize clone.
- We might inline f.

Where do such opportunities arise? (i) calling library code (ii) object-oriented programs.

Facts about interprocedural dataflow analysis

- Significantly harder asymptotically and to implement than intraprocedural dataflow analysis
 - Intraprocedural analysis: unknowns are lattice values.
 - Interprocedural analysis: unknowns are functions on lattice values.
 - **Aliasing**: different program names for same location
- Strategies for dealing with complexity:
 - invent special-purpose algorithms that work for important special cases (eg, if the lattice is finite or of bounded height)
 - use general-purpose techniques that compute approximate but conservative solutions

Game plan for inter-procedural analysis lectures:

- Start with call-by-value language
Key problem: solving dataflow function equations
- Call-by-value + global variables
Some problems have structure we can exploit to speed up analysis.
- Call-by-reference language
Additional problem: **aliasing**

Let us begin with simple program model in which there is no aliasing

```
MAIN()
  var p,q;
  p = read();
  q = f(p,3);
  ...
PROCEDURE f(x,y)
  var z,a;
  if (x > 0) z = x;
  a = y;
  ...f(y,z)...
```

- no higher-order procedures
- no data structures
- no global variables
- call-by-value
- assignments in procedure modify locals and parameters
- recursion is allowed

Check: no aliasing

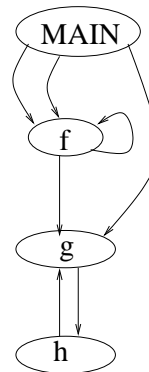
Key data structure: Call (multi)graph

- Structure:

Nodes: one for each procedure

Edges: from node f to node g if procedure f may invoke procedure g

```
MAIN ()  
  ... f(...)  
  ... g(...)  
  ...f(...)  
  ...  
f(..)  
  ...f(...)  
  ...g(...)  
  
g(..)  
  ...h(...)..  
  
h(..)  
  ...g(...)...  
  
Program
```



Call Graph

- **Algorithm:** Building call graph is trivial if there are no higher-order procedures.
- **Use:** Call graph plays a role sort-of like that of control flow graph in intraprocedural analysis, but not quite....

Context-insensitive analysis

One obvious approach: reduce interproc case to intraproc case

forward dataflow problem:

- merge information from all call sites of procedure at START
- copy dataflow information coming out of END to all return sites

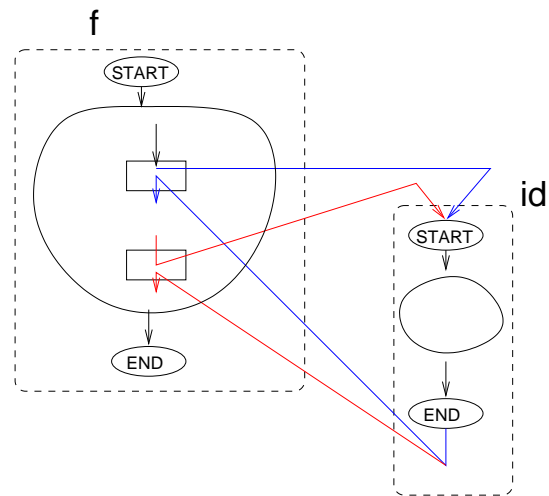
PROCEDURE f (..)

..... id(2)

.....id(3).....

PROCEDURE id(n)

return n;



Problem: information propagates along impossible interprocedural paths
such as blue edge into id and red edge out of id

This loses precision: in our example, we would not detect that id(2) is 2!

However, this reduction of inter-procedural analysis to the intra-procedural case
is safe.

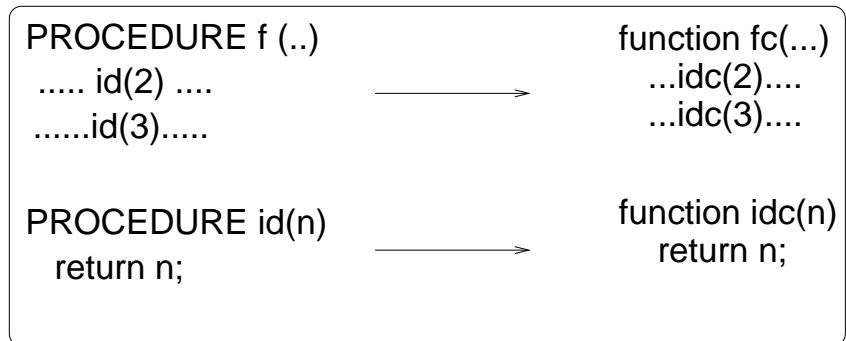
It is called context-insensitive analysis.

Context-sensitive analysis

Do not mix dataflow information from different call-sites of a procedure

One implementation of context-sensitive analysis:

- model each program procedure by a function on dataflow values
- to analyze dataflow effect of a call to procedure g , dataflow analyzer invokes the associated function, passing it some dataflow values and getting dataflow values back
- solves problem of avoiding dataflow mixing



Dataflow functions for constant propagation

Another implementation: build the composite graph as in context-insensitive analysis but propagate dataflow values together with a "tag" that identifies call sequence that generated that value (Sharir and Pnueli)

Main problem with context-sensitive analysis: termination

- Intuitively, we are doing something similar to a symbolic execution of program.
- Analysis must terminate even if program execution does not terminate!
- Difficulty with recursive procedures: analysis usually requires symbolic execution of both sides of conditionals, so how do we ensure termination? Here's an example where program terminates but analysis would not.

```
procedure main() {  
    var p;  
    p = read();  
    return f(3,p);  
}
```

```
procedure f(n,p) {  
  
    if (n > p)  
        then return f(n-1,p);  
        else return 1;  
}
```

```
function mainc() {  
    p = bottom;  
    p = T;  
    return fc(3,p);  
}
```

```
function fc(n,p) {  
    t1 = t2 = t3 = bottom;  
    if (n > p)  
        then t1 = fc(n-1,p);  
        else t2 = 1  
    t3 = join(t1,t2);  
    return t3;  
}
```

Source of termination problem:

The recursive definition of `fc` in previous slide is really an equational definition of `fc`: interpreting this definition as an “executable function” gets us into trouble in general.

Two important special cases:

- **No recursion in program**: no problem with non-termination in interpreting equational definitions as functions.

Determining there is no recursion: call-graph should have no cycles.

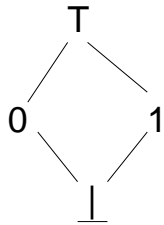
This idea does not work even in the presence of *static* recursion which even FORTRAN allows. **Static recursion**: text of program has recursive calls; **Dynamic recursion**: at runtime, two or more activations of a procedure coexist at some point in time.

- **Domain is finite**: solve equations iteratively by tabulating values of functions.

Solving recursive function equations by tabulation

- necessary condition: finite lattice

- Example: constant propagation in which all values other than 0 and 1 are set to T.



```
function fc(n,p)
  {t1 = t2 = t3 = bottom;
   if (n>p) then t1 = fc(n-1,p);
               else t2 = 1;
   t3 = join(t1,t2);
   return t3;}
```

Compute a sequence of approximations to fc as follows:

$$fc[0](n,p) = \perp$$

```
fc[i+1](n,p) =
  {t1 = t2 = t3 = bottom;
   if (n>p) then t1 = fc[i](n-1,p);
               else t2 = 1;
   t3 = join(t1,t2);
   return t3;}
```

For our example:

$$fc(n,p) = 1$$

Computing each element of sequence: make a table of output for each possible input value

Checking convergence: same table is obtained for two successive elements in sequence.

Termination: from monotonicity and finiteness of domain.

What do we do if we have recursion and domain is not finite?

Usual strategy: Replace recursive dataflow function with non-recursive conservative approximation

1. Identify recursive calls in call graph, and approximate their return values to T. Then, solve resulting acyclic problem.

```
function fc(n,p){  
  t1 = t2 = t3 = bottom;  
  if (n > p)  
    then t1 = fc(n-1,p); --->  
    else t2 = 1;  
  t3 = join(t1,t2);  
  return t3;}
```

```
function fc(n,p) {  
  t1 = t2 = t3 = bottom;  
  if (n > p)  
    then t1 = T;  
    else t2 = 1  
  t3 = join(t1,t2);  
  return t3;}
```

Recursive call identification: back edges in DFS of call-graph (not necessarily unique)

2. Solve context-insensitive problem and use result to approximate the effect of recursive calls.

- value on "return edge" will be an upper bound of possible return values
- replace recursive call with that value and solve acyclic context-sensitive problem

```
fc(n,p) {
```

```
  t1 = t2 = t3 = bottom;
```

```
  if (n > p)
```

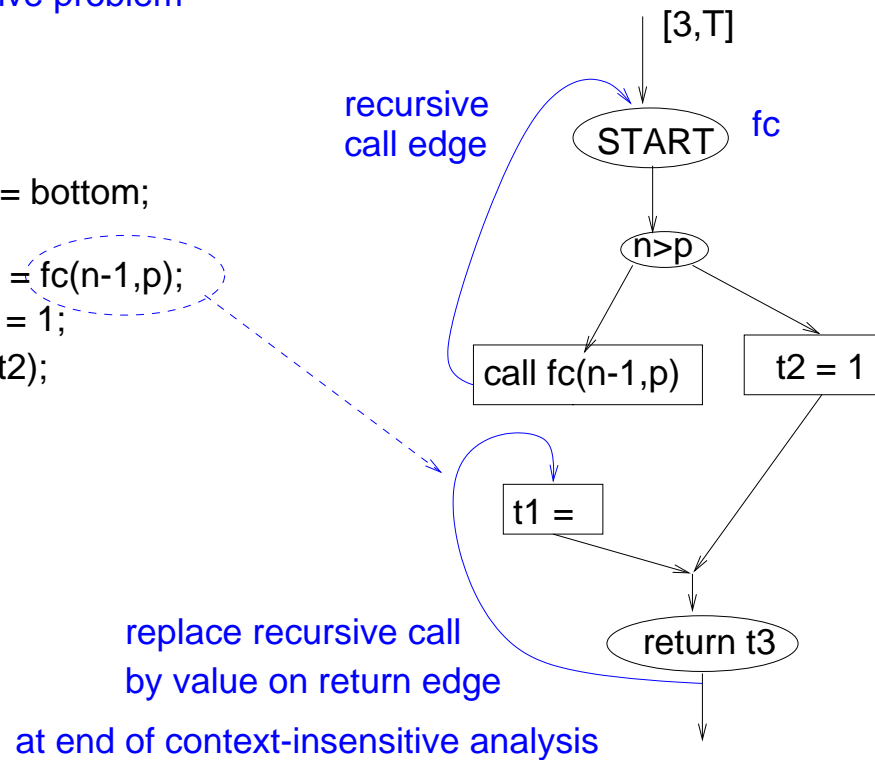
```
    then t1 = fc(n-1,p);
```

```
    else t2 = 1;
```

```
  t3 = join(t1,t2);
```

```
  return t3;
```

```
}
```



Adding global variables to program model:

```
GLOBAL G1, G2, G3
MAIN()
  var p,q
  p = f(G1,3); //S1 => MOD-S1 is {p,G2}
  ...
PROCEDURE f(x,y)
  var z,a
  if (x > 0) G2 := x;      // S2 => MOD-S2 is {G2}
  z = G1;                  // S3 => MOD-S3 is {z}
  ...f(G1,z)...
```

Still no aliasing.

Inter-procedural dataflow analysis: simple extension of call-by-value case.

(eg) Context-sensitive inter-procedural constant propagation: dataflow function for procedure will have one additional parameter for each global variable.

Exploiting structure in inter-procedural dataflow analysis

- Just as in intra-procedural case, inter-procedural problems may have structure that can be exploited to speed up solution.
- Exploiting intra-procedural structure: as before
- Inter-procedural structure: in the call graph
- For many problems, we can exploit strongly connected components in call graph to speed up analysis (eg. MOD computation)

Interprocedural dataflow problem: **computing MOD** [Banning]

*For any statement s , the set $MOD-s$ is the set of variables visible to s that **may** be modified directly or indirectly by execution of s .*

```
GLOBAL G1, G2, G3
MAIN()
  var p,q
  p = f(G1,3); //S1 => MOD-S1 is {p,G2}
  ...
PROCEDURE f(x,y)
  var z,a
  if (x > 0) G2 := x;    // S2 => MOD-S2 is {G2}
  z = G1;               // S3 => MOD-S3 is {z}
  ...f(G1,z)...
```

Auxiliary sets: For any procedure f , $GMOD-f$ is the set of global variables that may be modified directly or indirectly by invoking f .

In example, $GMOD-MAIN = \{G2\}$, and $GMOD-f = \{G2\}$

For any statement s , $MOD-s$ is the union of

- set of variables that may be modified directly in statement (IMOD)
- set of globals that may be modified directly or indirectly by procedure invocations in s (GMOD)

So given $GMOD$ sets, MOD sets are easy to compute.

How do we compute GMOD sets?

Write down a set of lattice equations and solve them.

- Lattice: power-set of global variables
- Equations: if procedure f has assignments to globals G_i, G_j, \dots and it may invoke procedures g, h, \dots equation for GMOD- f is

$$\text{GMOD-}f = \{G_i, G_j, \dots\} \cup \text{GMOD-}g \cup \text{GMOD-}h \dots$$

GLOBAL G1, G2, G3;

MAIN ()

... f(...)

.... g(...)

...f(...)

...

f(..)

G1 := ...

...f(...)

...g(...)

g(..)

...h(...)..

G1 := ...

h(..)

G3 := ...

...g(...)..

GMOD-main = GMOD-f U GMOD-g

GMOD-f = {G1} U GMOD-g U GMOD-f

GMOD-g = {G1} U GMOD-h

GMOD-h = {G3} U GMOD-g

Program

Observations

- We can use any iterative technique we discussed in intra-procedural case to solve these inter-procedural equations.
- Is there structure that can be exploited to reduce number of iterations? Yes!!
 - In our problem, information flows from invoked procedure to invokee.
 - So consider equations in “reverse invocation order”. See next slide.

GLOBAL G1, G2, G3;

MAIN ()

... f(...)

.... g(...)

...f(...)

...

f(..)

G1 := ...

...f(...)

...g(...)

g(..)

...h(...)..

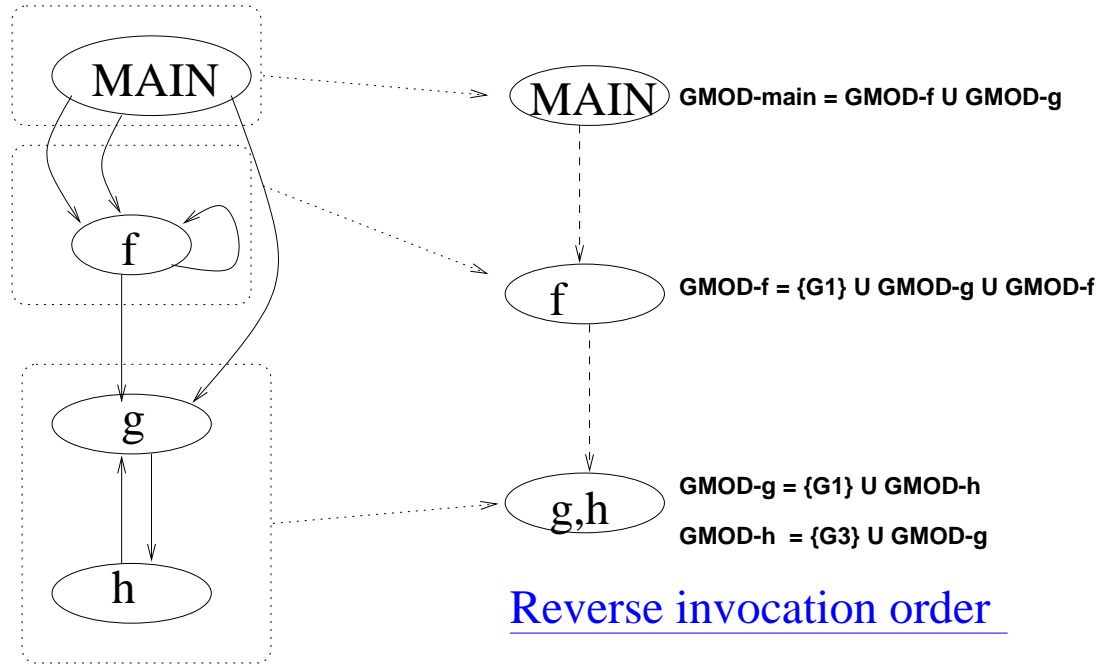
G1 := ...

h(..)

G3 := ...

...g(...)..

Program



Call Graph

Reverse invocation order

Further simplification: note that GMOD sets for all procedures in a single scc of call graph must be identical.

So collapse equations for all procedures in a single scc into a single equation!

$$\text{GMOD-g} = \{G1\} \cup \text{GMOD-h} \Rightarrow \text{GMOD-gh} = \{G1, G3\} \cup \text{GMOD-gh}$$

$$\text{GMOD-h} = \{G3\} \cup \text{GMOD-g}$$

Solving single equation for least solution: trivial! Just drop the recursive term.

$$\text{So in example, } \text{GMOD-g} = \text{GMOD-h} = \{G1, G3\}$$

Summary: GMOD/MOD computation for call-by-value language

- Write down GMOD equations for the program.
- Partition equations by scc's in call graph.
- Collapse equations in each scc into a single equation.
- In reverse topological order of acyclic condensate of call graph, read off solutions to GMOD equations.
- For each statement, compute MOD set.

Complexity: $O(\text{size of program} * \text{number of variables})$

Note: we can exploit scc's any time we have a set of equations (eg, block triangular systems in linear algebra)

Running example:

GLOBAL G1, G2, G3;

MAIN ()

... f(...)

... g(...)

...f(...)

...

f(..)

G1 := ...

...f(...)

...g(...)

g(..)

...h(...)..

G1 := ...

h(..)

G3 := ...

...g(...)..

Program

GMOD-main = GMOD-f U GMOD-g

GMOD-f = {G1} U GMOD-g U GMOD-f

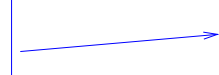
GMOD-g = {G1} U GMOD-h

GMOD-h = {G3} U GMOD-g

GMOD-main = {G1,G3}

GMOD-f = {G1,G3}

GMOD-gh = {G1,G3}



Call-by-reference

Complications:

1. Effect of a procedure is not just globals it modifies but also what happens to parameters

`x := f(G1,w) //to compute MOD, we need to know what happens to G1 and w!`

2. **Aliasing:** two program names for same location

```
GLOBAL G1,G2
procedure f(x,y)
  x := 3; //S1
  ....
procedure g(z,w)
  ..f(G1,G2)...f(z,z)...f(w,z)
```

For first call to f, x and y are not aliases

For second call to f, x and y are aliases

For third call to f, x and y are aliases if w and z are aliases!

What is MOD-S1???

Let us handle these problems one at a time.

Aliasing

- **MUST-ALIAS**: two program names that definitely refer to the same memory location.

```
GLOBAL G1;
```

```
procedure f(x) { <---- x and G1 are MUST-ALIASES within f  
  x = 7; }
```

```
procedure main() {  
  f(G1);  
  print(G1); }
```

MUST-ALIAS is an equivalence relation on names.

- **MAY-ALIAS**: two program names that may or may not refer to the same memory location. MAY-ALIASing usually arises from MUST-ALIASes through loss of information such as when we merge information along different program paths.

```
GLOBAL G1,G2;
procedure f(x) { <---- x and G1 are MAY-ALIAS's within f
    x = 7;}
procedure main() {
    f(G1) + f(G2);
    print(G1); }
```

MAY-ALIAS relation is reflexive and symmetric but not necessarily transitive.

Representation of aliasing information: alias pairs

```
GLOBAL G1,G2;
procedure f(x) { //ALIAS = {<x,x>,<x,G1>,<x,G2>}
  x = 7;}
procedure main() {
  f(G1) + f(G2);
  print(G1); }
```

If $\langle a,b \rangle$ does not occur in an ALIAS relation, variables a and b are definitely not aliased at the point in program where ALIAS relation holds.

Some people also store **may/must** flag with each alias pair.

Alias pairs are an example of **store-free alias representation**.

Store-based alias representation: see when we talk about pointer analysis.

Using ALIAS relations in inter-procedural dataflow analysis:

For our language model, all statements in a procedure have same alias relation (not true when we have pointers as in C).

Modify the dataflow transfer functions of statement with alias information

- Constant propagation:

```
procedure f(x,y) {      function fc(x,y) {
  ....                ....
  x := e;              V-out = {
                        let n = Eval(e,V-in);
                        return V-out where
                          V-out[i] = V-in[i] if i is not aliased with x
                          V-out[i] = join(Vin[i],n) if i is MAY-ALIASEd with x
                          V-out[i] = n if i is MUST-ALIASEd with x
                        }
  .....                .....
}                       }
```

- MOD computation: close affected variables under aliasing

```
GLOBAL G1,G2
procedure f(x,y)
  x := 3; // <--- MOD = {x,G1,y}
  ....
procedure g(z,w)
  ..f(G1,G2)...f(z,z)...f(w,z)
```

Computing alias relation for our program model: treat as a dataflow problem...

- Compute one alias set for each procedure.
- Each call-site has an associated transfer function that generates output alias set from alias set of caller (see next slide).
- Alias set of procedure = union of alias sets generated at its call sites.
- Transfer functions are monotonic and lattice is finite.


```

procedure f(x,y) {
  ...}
procedure g(a,b,c) { //suppose alias set is A
  ....f(p,q)... //output alias set is B
}

```

```

Transfer(A, list of actuals, list of formals, globals)
  B = { };
  for each actual parameter p in call do {
    if (p is a global variable) then append <p,x> to B;
    else for each tuple <p,V> in A do
      if V is a global variable then append <x,V> to B;
  }
  for each pair (p,q) of actual parameters in call do {
    if ((<p,q> is in A) or (p and q are same variable))
      then append <x,y> to B where x/y are formals bound to p/q by call.
  }
  return B;

```

Example: [adapted from Banning]

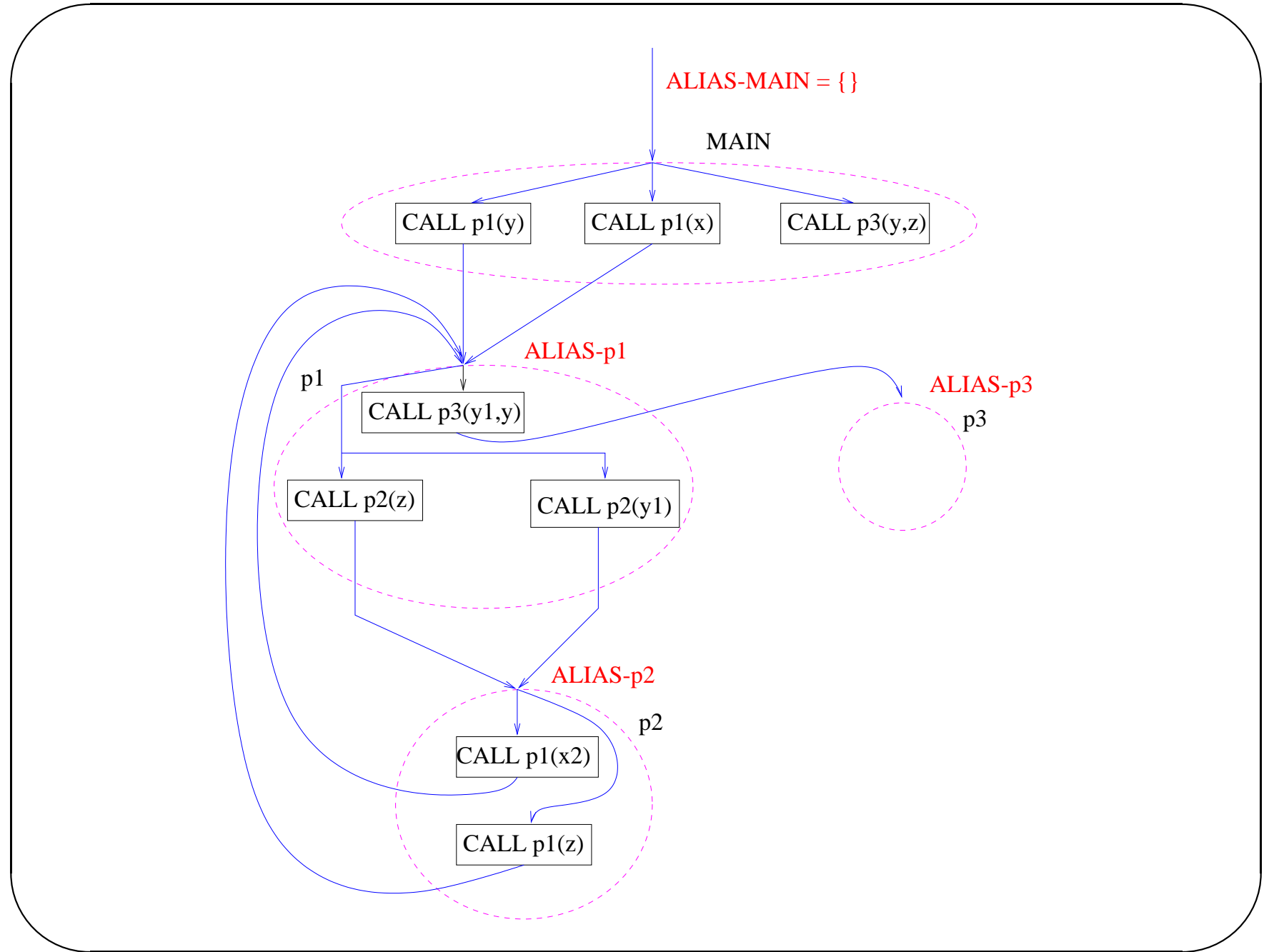
```
GLOBAL x,y,z;
procedure MAIN()      ALIAS-MAIN = {}
  p1(y);              ALIAS-C1  = {<y,y1>}
  p1(x);              ALIAS-C2  = {<x,y1>}
  p3(y,z);            ALIAS-C3  = {<y,x3>,<z,y3>}

procedure p3(x3,y3)  ALIAS-p3   = ALIAS-C3 U ALIAS-C6
  x3 := ...;
  y3 := ...;
  x  := ...;

procedure p1(y1)     ALIAS-p1   = ALIAS-C1 U ALIAS-C2 U ALIAS-C7 U ALIAS-C8
  p2(z);             ALIAS-C4 = {<x2,z>}
  p2(y1);            ALIAS-C5 = Transfer(ALIAS-p1, (y1)->(x2), {x,y,z})
  p3(y1,y);          ALIAS-C6 = Transfer(ALIAS-p1, (y1,y)->(x3,y3), {x,y,z})

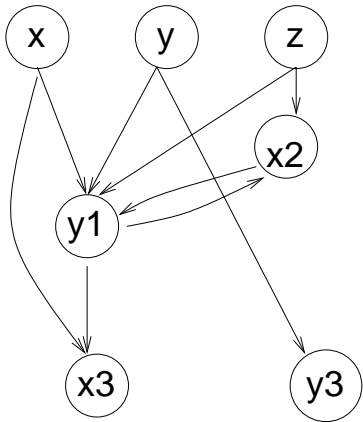
procedure p2(x2)     ALIAS-p2 = ALIAS-C4 U ALIAS-C5
  p1(x2);            ALIAS-C7 = Transfer(ALIAS-p2, (x2)->(y1), {x,y,z})
  p1(z);             ALIAS-C8 = {<z,y1>}

Solution: ALIAS-p1 = {<y,y1>,<x,y1>,<z,y1>}
ALIAS-p2 = {<y,x2>,<x,x2>,<z,x2>}
ALIAS-p3 = {<y,y3>,<x,x3>,<z,x3>,<y3,x3>,<y,x3>,<z,y3>}
```



More efficient ways of computing alias sets:

- Determine which globals are aliased with which formals by using [binding graph](#) (see next slide).
 - Graph has a node for each global and formal parameter; if global/formal v_1 is passed to formal v_2 , put an edge from v_1 to v_2 .
 - All formals reachable from node for global g are aliased to g .
- Determining which formals are aliased to each other: use [pairwise binding graph](#). Left to reader.



Binding graph

Structure of binding graph:

- one node for each reference parameter and global
- if procedure f passes its reference parameter/global r1 to procedure g as reference parameter r2, put an edge from r1 to r2

Finding aliases between globals and formals:

all reference parameters reachable from a global variable in the binding graph may be aliased to that global

In our example:

- x may be aliased to y1, x3, x2
- y may be aliased to y1, y3, x2,x3
- z may be aliased to y1, x2, x3

Concern 2:GMOD must tell us what happens to globals AND parameters.

One model: make GMOD into a function from variables to variables

```
w := f(G4,s); //S1
```

procedure f(x,y)	=>	GMOD-f(v1,v2)
var a,b;		return {G1,v1}
G1 := 5;		
if (y>x) x:= 7;		
...		

Intuition: (assuming s,w have no non-trivial aliases)

$$\text{MOD-S1} = \{w\} \cup \text{GMOD-f}(G4,s) = \{w\} \cup \{G1,G4\} = \{w,G1,G4\}$$

Example: [adapted from Banning]

```
GLOBAL x,y,z;
procedure MAIN()           GMOD-MAIN()
  p1(y);                   return GMOD-p1(y) U GMOD-1(x) U GMOD-p3(y,z)
  p1(x);
  p3(y,z);

procedure p3(x3,y3)       GMOD-p3(m,n)
  x3 := ...;              return {m,n,x}
  y3 := ...;
  x := ...;

procedure p1(y1)          GMOD-p1(m)
  p2(z);                  return GMOD-p2(z) U GMOD-p2(m) U GMOD-p3(m,y)
  p2(y1);
  p3(y1,y);

procedure p2(x2)          GMOD-p2(m)
  p1(x2);                 return GMOD-p1(m) U GMOD-p1(z)
  p1(z);
```

Need to solve recursive functional equations.

Without recursion, we can use “interpret equations as program” trick.

Another approach: since our lattice is finite, we can always use tabular approach.

GMOD-p3(m,n)

return {m,n,x}

GMOD-p1(m)

return GMOD-p2(z) U GMOD-p2(m) U GMOD-p3(m,y)

GMOD-p2(m)

return GMOD-p1(m) U GMOD-p1(z)

Iterations:

	1	2	3	... final
GMOD-p3(m,n):	{}	{m,n,x}	{m,n,x}	... {m,n,x}
GMOD-p2(m):	{}	{}	{}	... {m,y,x,z}
GMOD-p1(m):	{}	{}	{m,y,x}	... {m,y,x,z}

Exploiting structure for computing GMOD

- **call-by-value:** we found GMOD sets without iteration
 - find scc's in call graph
 - for each scc, compute union of side-effects to globals by procedures in scc
 - propagate GMOD sets in reverse invocation order
- can we use this trick for call-by-reference as well?
- **Problem:** GMOD sets contain both globals and parameters, so GMOD sets of mutually recursive procedures will be different in general.
- Idea: separate computation of side-effects to parameters from side-effects to globals

Side-effects to parameters:

procedure f(a,b) ...

$\text{RMODf-a} = \text{true}$ if execution of f modifies a directly or indirectly

```
GLOBAL x,y,z;
```

```
procedure MAIN()
```

```
  p1(y);
```

```
  p1(x);
```

```
  p3(y,z);
```

```
procedure p3(x3,y3)     $\text{RMODp3-x3} = \text{true}$ 
```

```
  x3 := ...;           $\text{RMODp3-y3} = \text{true}$ 
```

```
  y3 := ...;
```

```
  x := ...;
```

```
procedure p1(y1)       $\text{RMODp1-y1} = \text{RMODp2-x2} \vee \text{RMODp3-x3}$ 
```

```
  p2(z);
```

```
  p2(y1);
```

```
  p3(y1,y);
```

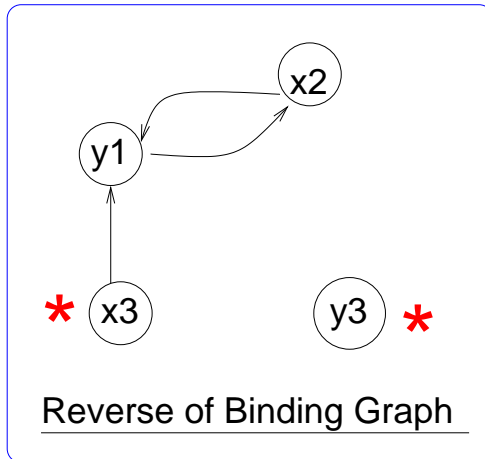
```
procedure p2(x2)       $\text{RMODp2-x2} = \text{RMODp1-y1}$ 
```

```
  p1(x2);
```

```
  p1(z);
```

Solution: $\text{RMODp3-x3} = \text{RMODp3-y3} = \text{RMODp1-y1} = \text{RMODp2-x2} = \text{true}$

Graphical way of solving RMOD equations: marker propagation



A reference parameter of a procedure f may be modified by execution of f if

- f may write to parameter directly
- f passes it to procedure g as a reference parameter, and g may modify the parameter

RMOD computation

- build reverse of binding graph (ignore globals)
- if f modifies reference parameter r1 directly, put a mark on r1
- propagate marks along reverse binding graph edges
- at the end of propagation, any variable that is marked corresponds to a reference parameter that may be modified

Side-effects to globals: similar to call-by-value

```
GLOBAL x,y,z;
procedure MAIN()
  p1(y);
  p1(x);
  p3(y,z);

procedure p3(x3,y3)    GLOBALp3 = {x}
  x3 := ...;
  y3 := ...;
  x  := ...;

procedure p1(y1)      GLOBALp1 = GLOBALp2 U GLOBALp3
  p2(z);
  p2(y1);
  p3(y1,y);

procedure p2(x2)      GLOBALp2 = GLOBALp1
  p1(x2);
  p1(z);
```

Solution: GLOBALp1 = GLOBALp2 = GLOBALp3 = {x}

From RMOD and GLOBALS, we can read off GMOD sets:

$$\text{GMODp1}(m) = \{x, m\}$$

$$\text{GMODp2}(m) = \{x, m\}$$

$$\text{GMODp3}(m, n) = \{x, m, n\}$$

Oops...

What went wrong?

We did not take into account side-effects to globals that were passed as parameters![Kennedy and Cooper]

Correct equations for GLOBAL sets: use RMOD information for globals passed as parameters:

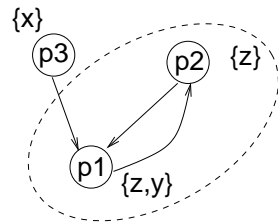
```
GLOBAL x,y,z;
procedure MAIN()
  p1(y);
  p1(x);
  p3(y,z);

procedure p3(x3,y3)    GLOBALp3 = {x}
  x3 := ...;
  y3 := ...;
  x  := ...;

procedure p1(y1)      GLOBALp1 = {z} U GLOBALp2 U GLOBALp3 U {y}
  p2(z);
  p2(y1);
  p3(y1,y);

procedure p2(x2)      GLOBALp2 = {z} U GLOBALp1
  p1(x2);
  p1(z);
```

This gives the correct sets.



Algorithm for side-effects to globals: GLOBAL

- Find acyclic condensate of the call graph.
- For each procedure, determine set of globals either assigned to directly in procedure or passed by reference as a parameter to a procedure that modifies that parameter (use RMOD information for this).
- Union these sets for all procedures in an scc.
- Propagate these global sets in reverse invocation order in the acyclic condensate.

Putting it all together: GMOD computation

- Compute RMOD information
 - build binding graph
 - mark every node that represents a reference parameter modified directly by its procedure
 - propagate marks in binding graph: efficient approach would compute scc's and propagate in acyclic condensate
 - for each procedure f , read off $\text{RMOD-}f =$ set of reference parameters of f that are marked.
- Compute GLOBALs information
 - build call graph
 - for each procedure, find all globals that are either modified directly by procedure or passed as a reference parameter to another procedure that modifies that parameter (use RMOD for this)

- find scc's of call graph and propagate sets in reverse invocation order.
- From RMOD and GLOBALs sets, read off GMOD function for each procedure f.

Summary

- Inter-procedural dataflow analysis: unknowns are functions on dataflow values
- Need to solve recursive functional equations
- Important special cases for which exact solution is possible
 - finite lattice: use tabular method
 - acyclic call graph: interpret equations as program
- Approximate solution of functional equations: approximate effect of backedges in call-graph
- Call-by-reference: need to take aliases into account
- Some interprocedural dataflow analysis problems can be reduced to marker propagation by formulating in the right graph. Key structure to exploit: strongly connected components in call graph.